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## (54) AMIDE COMPOUNDS AND USE OF THE SAME

(57) An amide compound of the formula (I):

$$R - A - X \xrightarrow{R^{1}} R^{2} \xrightarrow{0} (CH_{2})_{m} R^{6}$$

$$R - A - X \xrightarrow{R^{3}} R^{4} \qquad R^{5}$$
(1)

wherein R is amino and the like, A is alkylene and the like, X is O, S and the like, M is arylene and the like,  $R^1$ ,  $R^2$ ,  $R^3$  and  $R^4$  are H, hydroxy and the like,  $R^5$  is H, alkyl and the like, m is an integer of 0-6,  $R^6$  is an optionally substituted aryl and the like, and  $R^7$  is H, an optionally substituted alkyl and the like, a pharmaceutically acceptable acid addition salt thereof and a pharmaceutical containing same as an active ingredient. The amide compounds exhibit superior suppressive effects on cytokines directly or indirectly involved in inflammations, such as IL-8, IL-1, IL-6, TNF- $\alpha$ , GM-CSF and the like, and are useful for the prophylaxis and treatment of rheumatic diseases, arthritis due to gout and the like.

## Description

## Technical Field

The present invention relates to a novel compound exhibiting superior suppressive effects on cytokines directly or indirectly involved in inflammations, such as interleukin-8 (IL-8), interleukin-1 (IL-1), interleukin-6 (IL-6), tumor necrosis factor (TNF-α), GM-CSF and the like, and pharmaceutical agents comprising said compound, such as anti-inflammatory agents.

## Background Art

An inflammation is one of the protective responses in the living organisms which aims at removal of foreign substances, pathogenic bacteria and so on, as well as repair of damaged tissues. When inflammatory stimulation is received, the microcirculatory system responds and particularly increases vascular permeability. The vascular permeability is promoted by chemical mediators and cytokines. Sequentially, chemotaxis, migration and activation of neutrophiles are induced, foreign substances and pathogenic bacteria are phagocytosed at the sites of inflammation, and chemical mediators are released to induce inflammatory responses. Subsequent to neutrophiles, chemotaxis and recruitment of macrophages at the local sites occur, and activated macrophages, like neutrophiles, phagocytose foreign substances, pathogenic bacteria, disintegrated tissues and so on to produce various cytokines. Then, pathogenic bacteria, foreign substances and damaged tissues are removed and the tissues are re-constructed, whereby the inflammation comes to an end. The above-mentioned process occurs in normal inflammatory responses. In allergy and autoimmune diseases such as rheumatoid arthritis and systemic lupus erythematosus, however, abnormal immune responses prolong inflammation and cause strong systemic symptoms.

Many cytokines are involved in various processes of inflammatory responses. For example, IL-1, TNF- $\alpha$  and IL-8 are responsible for the chemotaxis, adhesion to vascular endothelial cells, and migration into vascular walls, of leukocytes, which are seen during migration of leukocytes into the sites of inflammation, wherein IL-1, TNF- $\alpha$  and IL-8 activate neutrophiles to cause release of lysosomal enzymes and production of active oxygen and prostaglandin, thus inducing inflammation. When IL-1, TNF- $\alpha$  and IL-6 migrate into the circulatory system, they act on liver to induce production of acute phase inflammatory protein (e.g., CRP and SAA), and act on bone marrow to increase neutrophiles and platelets. In inflammations of connective tissues, such as rheumatoid arthritis (RA), IL-1 and TNF- $\alpha$  are said to activate fibroblasts and osteoclastic cells and induce production of prostaglandin and collagenase [Mebio, 11 (2), 18-23, (1994)].

As stated in the foregoing, IL-1 and TNF- $\alpha$  play a central role in various aspects of inflammatory responses.

Meanwhile, IL-8 is produced not only by peripheral blood monocytes and tissue macrophages, but also by large granular lymphocytes (LGL) known as natural killer cells, T lymphocytes and various tissues and cells such as fibroblasts, vascular endothelial cells and epidermal keratinocytes. Examples of production stimulators include mitogen lectins such as LPS, PHA, PSK (Coriolus versicolor-derived protein-bound polysaccharide, Krestin) and cytokines such as IL-1 and TNF-α.

Although most of these cells barely produce IL-8 constantly, upon stimulation with the above-mentioned IL-8 production stimulators, they produce more than 100 times greater amounts of IL-8 within 24 hours as compared to the production without stimulation. For example, when human peripheral blood monocytes are stimulated with PSK, IL-8 mRNA is induced within an hour, and production amount of IL-8 mRNA reaches its peak in 3 hours, and gradually decreases with time. Along with the induction of IL-8 mRNA, IL-8 protein having neutrophile chemotaxisis ability is detected in the medium at 3 hours after the stimulation and increases with time. IL-8 mRNA is induced in the same manner in time as in the stimulation of IL-1 and TNF-α. IL-8 is noticeably stable to protease produced by activated macrophage and the like.

The in vitro biological activities of IL-8 include chemotactic promotion, induction of degranulation, respiratory burst induction, lysosomal enzyme release induction, induction of adhesion to unstimulated or stimulated vascular endothelial cells, promotion of extravascular migration, reinforcement of expression of adhesion factors, leukotriene B<sub>4</sub>-HETH release induction and the like with regard to neutrophiles; chemotactic promotion with regard to T cells; suppressive effect on IgE production by IL-4 with regard to B cells; and chemotactic promotion and histamine · leukotriene release induction with regard to basophils. IL-8 also has in vivo activities of induction of migration of neutrophiles and lymphocytes, induction of neutrophilia, reinforcement of vascular permeability, and neutrophile-dependent arthrosynovial destruction [Rinsho Men-eki, 25 (8), 1013-1020 (1993)].

As mentioned earlier, IL-8 has various effects on neutrophiles. It has been gradually clarified that IL-8 also acts on T lymphocytes, basophils, monocytes, keratinocytes and melanoma cells, besides neutrophiles. The biological activities and target cells thereof are found to be diverse like other cytokines.

It has been known that IL-8 realizes, in vivo, migration of neutrophiles and lymphocytes at the sites of subcutaneous

injections, and increases homing of T lymphocytes to local lymph nodes. It has been also known that an intravenous or intraperitoneal injection of IL-8 markedly increases neutrophile counts in peripheral blood, and administration in large amounts thereof causes destruction of alveoli. In addition, an injection of IL-8 into rabbit intra-articular joint space is known to lead to arthrosynovial destruction with migration of large amounts of neutrophiles. These results suggest strong inflammation induction by IL-8 in vivo.

In view of the fact that IL-8 has various actions besides chemotactic stimulation of neutrophile, that IL-8 was detected in synovial fluid in patients with gout or rheumatic arthritis, that IL-8 was detected from skin pieces of patients with dermatitis such as psoriasis, that IL-8-like chemotactic factor is produced by peripheral blood monocytes in asthma, and that IL-8 was detected in peripheral blood of patients with sepsis which is considered to be one of the causes of adult respiratory distress syndrome (ARDS), it is evident that IL-8 is involved in various diseases such as inflammation.

Therefore, a substance capable of suppressing cytokines responsible for inflammations, such as IL-1, IL-6, IL-8 and TNF- $\alpha$ , is extremely useful as a new type of medicine for noninfectious or infectious diseases accompanied by neutrophile migration, which are represented by rheumatic diseases (e.g., rheumatoid arthritis); arthritis due to gout; systemic lupus erythematosus; dermatopathy (e.g., psoriasis, pustulosis and atopic dermatitis); respiratory diseases (e.g., bronchial asthma, bronchitis, ARDS and diffused interstitial pneumonia); inflammatory bowel diseases (e.g., ulcerative colitis and Crohn's disease); acute or chronic hepatitis inclusive of fulminant hepatitis; acute or chronic glomerulone-phritis; nephropyelitis; uveitis caused by Behcet disease and vogt-Koyanagi Harada disease; Mediterranean fever (polyserositis); ischemic diseases (e.g., myocardial infarction); and systemic circulatory failure and multi-organ failure caused by sepsis. In particular, such substance is expected to be effective as an anti-inflammatory agent based on new action mechanisms.

With such background of the art, compounds having inhibitory activity on inflammatory cytokines, such as IL-8, have been recently reported. For example, Japanese Patent Application under PCT laid-open under Kohyo No. 7-503017 discloses an imidazole derivative such as 4-(4-fluorophenyl)-2-(4-methylthiophenyl)-5-(4-pyridyl)imidazol as a cytokine inhibitor; Japanese Patent Application under PCT laid-open under Kohyo No. 7-503018 discloses pyridyl-substituted imidazole derivatives such as 1-(4-pyridyl)-2-(4-fluorophenyl)-4-phenylimidazol as cytokine inhibitors; and Japanese Patent Unexamined Publication No. 3-34959 discloses naphthalenemethaneamino derivatives having cytokine inhibitory activity. However, these publications do not suggest the compound of the present invention.

In addition, compounds having inhibitory activity on protease involved in inflammatory diseases have been reported. For example, Japanese Patent Unexamined Publication No. 4-330094 discloses t-butyloxycarbonyl-trimethyl-silyl-Ala-Pro-NH-CH[(CH<sub>2</sub>)<sub>3</sub>N<sub>3</sub>]-B-pinandiole as a serine protease inhibitor of thrombin which induces pre-inflammatory changes of IL-1 and the like. Japanese Patent Examined Publication No. 7-53705 discloses phenylalanine derivatives such as N-(trans-4-aminomethylcyclohexylcarbonyl)-L-phenylalanine 4-acetylanilide. However, this publication relates to a compound characteristically having amino at one end of phenylalanine and 4-aminomethyl-6-membered ring-carbonyl group at the other end, which relates to a protease inhibitor, and does not relate to an inflammatory cytokine production suppressor, such as the compound of the present invention.

An object of the present invention is to provide a compound usable as a novel selective anti-inflammatory agent which suppresses production and release of inflammatory cytokines such as IL-8, IL-1, TNF-α, IL-6, and the like.

In addition, an object of the present invention is to provide a pharmaceutical agent comprising said compound.

## Disclosure of the Invention

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The present inventors have conducted intensive studies with the aim of achieving the above-mentioned objects and completed the present invention.

Accordingly, the present invention provides the following.

# (1) An amide compound of the formula (1):

$$R - A - X \xrightarrow{R^1 \qquad R^2 \qquad 0} \bigvee_{R^3 \qquad R^4 \qquad R^5} (CH_2)_{\infty}^{R^6}$$

$$(I)$$

wherein;

	R	is an optionally substituted non-aromatic heterocyclic group containing nitrogen, a hydroxy, $R_a$ , an alkoxy substituted by $R_a$ , an alkylthio substituted by $R_a$ , or an alkylamino substituted by $R_a$ .
5		wherein $R_a$ is amino, guanidino, amidino, carbamoyl, ureido, thioureido, hydrazino, hydrazino-carbonyl or imino, these groups being optionally substituted by a substituent selected from the group consisting of lower alkyl, halogenated lower alkyl, cycloalkyl, aralkyl, aryl and amino-protecting group;
f	Α	is an optionally substituted, linear or branched alkylene which optionally has one or more double bond(s) or triple bond(s) in the chain, or a single bond;
10	X	is an oxygen atom, a sulfur atom, a cycloalkylene, a divalent aromatic heterocyclic group having one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, -SO-, -SO <sub>2</sub> -, -C=C-, -CO-, -CO-, -CO-, -CO-, -CS-, -CO-CO-0-, -NH-CO-NH-, -NH-CS-NH-, -NH-C(=NH)-NH-, -NR <sup>8</sup> -, -NR <sup>8</sup> CO-, -CONR <sup>8</sup> -, -NR <sup>8</sup> SO <sub>2</sub> -, -SO <sub>2</sub> NR <sup>8</sup> -, -NR <sup>8</sup> -COO-, -OOC-NR <sup>8</sup> -, or -CR <sup>9</sup> R <sup>10</sup> -
15	M	wherein R <sup>8</sup> is hydrogen atom, alkyl, cycloalkyl, aryl, aralkyl or amino-protecting group, and R <sup>9</sup> and R <sup>10</sup> are the same or different and each is hydrogen atom, alkyl, cycloalkyl, aryl or aralkyl; is an arylene, a cycloalkylene, or a divalent heterocyclic group which has one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom,
20	R <sup>1</sup> , R <sup>2</sup> , R <sup>3</sup> and R <sup>4</sup>	and which optionally forms a fused ring; are the same or different and each is a hydrogen atom, a hydroxy, a halogen atom, an alkoxy, a mercapto, an alkylthio, a nitro, a cyano, a carboxy, an alkoxycarbonyl, an acyl, an alkyl optionally substituted by a substituent selected from the group consisting of hydroxy, lower alkoxy and halogen atom, an amino optionally substituted by a substituent
25		selected from the group consisting of alkyl, aryl, aralkyl and amino-protecting group, or -O-CO-R <sup>11</sup> wherein R <sup>11</sup> is optionally substituted alkoxy, optionally substituted aryl, optionally substituted
30		cycloalkyl, optionally substituted aryloxy, optionally substituted aralkyloxy, optionally substituted alkylthio, optionally substituted arylthio, or alkyl optionally substituted by a substituent selected from the group consisting of alkoxycarbonyl, acyloxy, aryl, aryloxy, aryloxycarbonyl, aralkyloxy, aralkyloxycarbonyl, alkylthio, arylthio, acyl, lower alkoxy, carboxy, halogen atom
	R <sup>5</sup>	and amino optionally substituted by lower alkyl or acyl; is a hydrogen atom, an alkyl optionally substituted by halogen atom, an optionally substituted aralkyl, or an amino-protecting group; is 0 or an integer of 1-6;
35	m R <sup>6</sup>	is an optionally substituted aryl, an optionally substituted cycloalkyl, an optionally substituted lower alkyl, an optionally substituted lower alkoxy, an optionally substituted lower alkylthio, an amino optionally substituted by a substituent selected from the group consisting of lower alkyl, aryl, aralkyl and amino-protecting group, or an optionally substituted heterocyclic group having one ore more hetero atoms selected from the group consisting of a nitrogen atom, sulfur
40	R <sup>7</sup>	atom and oxygen atom; and is a hydrogen atom, an optionally substituted alkyl, an optionally substituted aryl, an optionally substituted aromatic heterocyclic group having one ore more hetero atoms selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, or -CO(Y) <sub>p</sub> R <sup>12</sup> wherein Y is oxygen atom, sulfur atom, -NR <sup>13</sup> - or -NR <sup>13</sup> -SO <sub>2</sub> -wherein R <sup>13</sup> is hydrogen atom,
45		alkyl, aralkyl, hydroxy, alkoxy, aryl or amino-protecting group, p is 0 or 1, and R <sup>12</sup> is hydrogen atom, optionally substituted alkenyl, optionally substituted alkynyl, optionally substituted cycloalkyl, optionally substituted aryl, optionally substituted aralkyl, adamantyl, cycloalkylideneamino, optionally substituted heterocyclic group having one or more hetero atom(s) selected from the group consisting of nitrogen atom, sulfur atom and oxygen atom, or alkyl
50		optionally substituted by a substituent selected from the group consisting of hydroxy, alkoxy, alkoxyalkoxy, alkoxycarbonyl, acyloxy, carboxy, heterocyclic group having one or more hetero atom(s) selected from the group consisting of nitrogen atom, sulfur atom and oxygen atom, and amino optionally substituted by a substituent selected from the group consisting of alkyl, aryl, aralkyl and amino-protecting group;
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and a pharmaceutically acceptable acid addition salt thereof. (2) The amide compound of (1) above, wherein, in the formula (I), at least one symbol selected from the group consisting of R, A, X, M,  $R^1$ ,  $R^2$ ,  $R^3$ ,  $R^4$ ,  $R^5$ , m,  $R^6$  and  $R^7$  satisfies the following definitions, and a pharmaceutically

	or and thereof:
acceptable acid addi	tion salt thereof:  is a non-aromatic heterocyclic group containing nitrogen, which is optionally substituted by  is a non-aromatic heterocyclic group, Ra1, an alkoxy substituted by Ra1, an alkylthio substi-
	is a non-aromatic heterocyclic group containing nitrogen, which is optionally substituted by R <sub>a1</sub> , an alkylthio substituted by R <sub>a1</sub> , an alkylthio substituted by R <sub>a1</sub> , an alkylamino substituted by R <sub>a1</sub> ,
R	lower alkyl of all life production and the state of the s
	tuted by R <sub>a1</sub> , or an alkylamino substituted by R <sub>a1</sub> , tuted by R <sub>a1</sub> , or an alkylamino, amidino, carbamoyl, ureido, thioureido, hydrazino, hydrazino by R <sub>a1</sub> , or an alkylamino, amidino, carbamoyl, ureido, thioureido, hydrazino hydrazino by a substituent selected from
5	tuted by R <sub>a1</sub> , or an alkylamino substituted by a substituent selected from wherein R <sub>a1</sub> is amino, guanidino, amidino, carbamoyl, ureido, thioureido, nydrazino, nydrazino, substituent selected from nocarbonyl or imino, these groups being optionally substituted by a substituent selected from nocarbonyl or imino, these groups being optionally substituted by a substitutent selected from nocarbonyl or imino, these groups being optionally substituted by a substituted from nocarbonyl or imino, these groups being optionally substituted by a substitut
	nocarbonyl or imino, these groups being optionally substitutes of course in the group consisting of lower alkyl, aralkyl and amino-protecting group; the group consisting of lower alkylene which optionally has one or more double bond(s) or triple
	the group consisting of the dense which potionally has one of the constant and the dense which potionally has one of the constant and the cons
Α	is a linear or branched alkylerie which specified in the chain, or a single bond; bond(s) in the chain, or a single bond; bond(s) in the chain, a sulfur atom, a cycloalkylene, a divalent aromatic heterocyclic group havis an oxygen atom, a sulfur atom, a cycloalkylene, a divalent aromatic heterocyclic group havis an oxygen atom, a sulfur atom, a cycloalkylene, a divalent aromatic heterocyclic group havis an oxygen atom, a sulfur atom, a cycloalkylene, a divalent aromatic heterocyclic group havis an oxygen atom, a sulfur atom, a cycloalkylene, a divalent aromatic heterocyclic group havis an oxygen atom, a sulfur atom, a cycloalkylene, a divalent aromatic heterocyclic group havis an oxygen atom, a sulfur atom, a cycloalkylene, a divalent aromatic heterocyclic group havis an oxygen atom, a sulfur atom, a cycloalkylene, a cycloalkyl
10	bond(s) in the chain, or a single bond, bond(s) in the chain, or a single bond, is an oxygen atom, a sulfur atom, a cycloalkylene, a divalent aromatic neterocyclic group is an oxygen atom, a sulfur atom, a cycloalkylene, a divalent aromatic neterocyclic group is sufficient atom, a cycloalkylene, a divalent aromatic neterocyclic group, sulfur is an oxygen atom, a cycloalkylene, a divalent aromatic neterocyclic group, sulfur is an oxygen atom, a cycloalkylene, a divalent aromatic neterocyclic group, sulfur is an oxygen atom, a cycloalkylene, a divalent aromatic neterocyclic group, sulfur is an oxygen atom, a cycloalkylene, a divalent aromatic neterocyclic group, sulfur is an oxygen atom, a cycloalkylene, a divalent aromatic neterocyclic group, sulfur is an oxygen atom, a sulfur atom, a cycloalkylene, a divalent aromatic neterocyclic group, and the cycloalkylene, a cycloalkylene, a divalent aromatic neterocyclic group, and the cycloalkylene, a cycl
· X	ing one or more hetero atom(s) selected from the group and one or more hetero atom(s) selected from the group and one or more hetero atom(s) selected from the group and one or more hetero atom(s) selected from the group and one or more hetero atom(s) selected from the group and one or more hetero atom(s) selected from the group and one or more hetero atom(s) selected from the group and one or more hetero atom(s) selected from the group and one or more hetero atom(s) selected from the group and one or more hetero atom(s) selected from the group and one or more hetero atom(s) selected from the group and one or more hetero atom(s) selected from the group and one or more hetero atom(s) selected from the group and one or more hetero atom(s) selected from the group and one or more hetero atom(s) selected from the group and one or more hetero atom(s) selected from the group and one or more hetero atom(s) selected from the group and one or more hetero atom(s) selected from the group and one or more hetero atom(s) selected from the group and one or more than the group and one o
	atom and oxygen atom, and All NH-C(=NH)-NH-, -INT
	CO-O-, -NH-CO-NH-, -NH-CO-NH-, -NH-COO-, -OOC-NR <sup>8</sup> -, or -CR <sup>9-R 10-</sup>
_	NR°SO <sub>2</sub> -, -SO <sub>2</sub> Nr1   NR°SO <sub>2</sub> Nr1   NR°SO <sub>2</sub> Nr2   NR°SO <sub>2</sub>
15	wherein House is hydrogen atom, lower arry or more hetero
	are the same or division and division or a division thereocyclic group must atom and oxygen atom,
M	are the same or different and each is hydrogen are the same or different and each is hydrogen which has one or interest is an arylene, a cycloalkylene or a divalent heterocyclic group which has one or with a strong and oxygen atom, is an arylene, a cycloalkylene or a divalent heterocyclic group which has one or different and each is hydrogen atom, and oxygen atom, a lower atom of the strong is a strong in the strong is a strong in the strong in the strong in the strong is a strong in the strong in the strong in the strong is a strong in the s
•	atom(s) selected from the group consisting and which optionally forms a fused ring; and which optionally forms a fused ring; are the same or different and each is a hydrogen atom, a hydroxy, a lower alkoxycarbonyl, an are the same or different and each is a hydrogen atom, a carboxy, a lower alkoxycarbonyl, an are the same or different and each is a hydrogen atom, a chapter alkoxycarbonyl, an are the same or different and each is a hydrogen atom, a hydroxy, a halogen atom, a lower alkytthio, a nitro, a cyano, a carboxy, a lower alkoxycarbonyl, an
20 R <sup>1</sup> , R <sup>2</sup> , R <sup>3</sup> and F	and which optionally forms a tused might and which optionally forms a tused might and which optionally forms a tused might and which optionally a hydroxy, a hard and each is a hydrogen atom, a hydroxy, a lower alkoxycarbonyl, an are the same or different and each is a hydrogen atom, a carboxy, a lower alkylthio, a nitro, a cyano, a carboxy, a lower alkylthio, a nitro, a cyano, a carboxy, a lower alkylthio, a nitro, a cyano, a carboxy, a lower alkylthio, a nitro, a cyano, a carboxy, a hydroxylthio optionally substituted by a substituted substituted
R', H-, h and	alkoxy, a mercapio, a total larger alkyl optionally substituted by a substituted
	are the same or different and each is a without a cyano, a carboxy, a lower alkoxyocarbonyl alkoxy, a mercapto, a lower alkylthio, a nitro, a cyano, a carboxy, a lower alkoxyocarbonyl, an acyl, a lower alkyl optionally substituted by a substituent selected from the group consisting of hydroxy, lower alkoxy and halogen atom, an amino optionally substituted group consisting of lower alkyl, aralkyl and amino-protect-by a substituent selected from the group consisting of lower alkyl optionally substituents.
	group consisting of flyer who group consisting of lower airly, all airly
25	by a substituted cycloalkyl, lower alkyl optionally substi-
	by a substituent selected from the group established by a substituted cycloalkyl, lower alkyl optionally substituted cycloalkyl, lower alkyl optionally substituted cycloalkyl, lower alkoxycarbonyl, acyloxy, wherein R <sup>11</sup> ' is lower alkoxy, optionally substituted by tuted by a substituent selected from the group consisting optionally substituted by a substituted by a substituted by a substituted by a substituted by the substitute
	tuted by a substituent selected from the group carboxy and amino optionally substituted by
	aralkyloxy, araikyloxycanists, heatinged by a substituent selection in the
	tuted by a substituent selected from the group consisting aralkyloxy, aralkyloxycarbonyl, acyl, lower alkoxy, carboxy and amino optionally substituted aralkyloxy, aralkyloytonally substituted by a substituent selected from the group consisting lower alkyl, or aryl optionally substituted by a substituted by halogen atom, an optionally substituted by halogen atom, an optionally substituted by halogen atom.
30	of lower alkyl, carbox, an alkyl optionally substituted by naroger around
R <sup>5</sup>	is a hydrogen atom, co-protecting group; aralkyl, or an amino-protecting group;
	a and addr of the
m	is 0 or an integer of vivial states of the s
<sub>35</sub> R <sup>6</sup>	from the group consisting and heterocyclic group having one atom are option-
	wherein said aryl, cycloalkyl and neterous memory alternation and oxygen atom and oxygen atom selected from the group consisting of nitrogen atom, sulfur atom and oxygen atom selected from the group consisting of lower alkyl, halogen ally substituted by a substituent selected from the group consisting of lower alkoxy, and lower alkoxycarbonyl; and
	selected from the group consisting
	ally substituted by a substituent selected from the group substituted by a substituted allower alkoxy, amino, carboxy and lower alkoxycarbonyl; and atom, hydroxy, lower alkoxy, amino, carboxy and lower alkoxycarboxy, lower alkoxycarboxy, lower alkoxycarboxy, lower alkoxycarboxy.
40	ally substituted by a substituent selected and lower alkoxycarbonyl; and atom, hydroxy, lower alkoxy, amino, carboxy and lower alkoxycarbonyl; and atom, hydroxy, lower alkoxy, amino, carboxy and lower alkoxycarbon, hydroxy, lower alkoxycarbon, lower alkylthio, carboxy, lower alkoxycarbon, lower alkylthio, carboxy, lower alkoxycarbon, lower alkylthio, carboxy, lower alkoxy, mercapto, lower alkylthio, carboxy, lower alkoxy, mercapto, lower alkylthio, carboxy, lower alkoxycarbon, lower alkoxycarbon, lower alkoxycarbon, lower alkoxycarbon, lower alkoxycarbon, lower alkoxycarbon, lower alkoxycarbonyl; and lower alkylthio, carboxy, lower alkylthio, carboxy, lower alkoxycarbonyl; and lower alkylthio, carboxycarbonyl; and lower alkylthio,
R <sup>7</sup>	atom, hydroxy, lower alkoxy, arinto, carboxy a substituted by a substituent selected that is a hydrogen atom, a lower alkyl optionally substituted by a substitutent selected that is a hydrogen atom, a lower alkyl optionally substituted by a substitutent selected that is a hydrogen atom, and substituted by a substitutent selected that is a hydrogen alkoxy, mercapto, lower alkylthio, carboxy, lower alkoxycar-group consisting of a nitrogen atom, sulfur atom and oxygen atom, and both is a hydrogen atom, and an around the group consisting of a nitrogen atom, sulfur atom and oxygen atom, and
	bonyl and arrillo, all all all and a nitrogen atom, suite all all all all all all all all all al
	selected from the group state allow or CO(Y)ph =13:- bustogen atom.
	which is optionally substitute of the NR <sup>13</sup> - or -NR <sup>13</sup>
<b>45</b> .	wherein Y is oxygen atom, alknow or amino-protecting groups it coult substituted by
	lower alkyl, atantyl, cycloalkylidenearillio, cycloalk
	lower alkyl, aralkyl, hydroxy, lower alkoxy of the lower alkyl, aralkyl, adamantyl, cycloalkylideneamino, cycloalkyl optionally substituted by a substituent selected from the group consisting of lower alkyl, alkyl optionally substituted by a substituent selected from the group consisting of, nitrohydroxy, lower alkoxy, low
	hydroxy, lower alroxy, toward better atom(s) selected from the substituent
50	rocyclic group riaving one atom, and amino optionally section group, aryl
•	gen atom, sumur alori and arising of lower alkyl, araikyl and arising alkyl, halo-
	selected from the group selected from the group constant which is ontionally
	optionally substituted by the standard lower alkoxy, or fleteroby the standard atom,
55	
	gen atom, amino, carboxy, hydroxy and lower alkyr, hardyes substituted by a substituent selected from the group consisting of lower alkyr, hardyes substituted by a substituent selected from the group consisting of nitrogen atom, and which has one or more hetero atom(s) selected amino, carboxy, hydroxy and lower alkoxy, and which has one or more hetero atom(s) selected amino, carboxy, hydroxy and lower alkoxy, and which has one or more hetero atom(s) selected amino, carboxy, hydroxy and lower alkoxy, and which has one or more hetero atom(s) selected amino, carboxy, hydroxy and lower alkoxy.
	amino, carboxy, hydroxy and lower alkoxy, and which has the or manner atom, from the group consisting of nitrogen atom, sulfur atom and oxygen atom.
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(3) The amide compound of (1) above, wherein, in the formula (I), at least one symbol selected from the group consisting of R, A, X, M,  $R^1$ ,  $R^2$ ,  $R^3$ ,  $R^4$ ,  $R^5$ , m,  $R^6$  and  $R^7$  satisfies the following definitions, and a pharmaceutically acceptable acid addition salt thereof:

is a non-aromatic heterocyclic group containing nitrogen, which is optionally substituted by

5	R	is a non-aromatic heterocyclic group containing nitrogen, which is optionally substituted by lower alkyl or amino-protecting group, $R_{a2}$ , or an alkoxy substituted by $R_{a2}$ , wherein $R_{a2}$ is amino, guanidino, amidino or carbamoyl, these groups being optionally substituted by lower alkyl or amino-protecting group;
	Α	is a linear alkylene or a single bond;
10	x	is an oxygen atom, a sulfur atom, a divalent aromatic heterocyclic group having one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, -COO-, -OOC-, -NR <sup>8</sup> ", -NR <sup>8</sup> "CO-, -CONR <sup>8</sup> "-, -NR <sup>8</sup> "SO <sub>2</sub> -, -SO <sub>2</sub> NR <sup>8</sup> "-, or -CR <sup>9</sup> "R <sup>10</sup> "- wherein R <sup>8</sup> " is hydrogen atom, lower alkyl or amino-protecting group, and R <sup>9</sup> " and R <sup>10</sup> " are the same or different and each is hydrogen atom or lower alkyl;
15	M	is an arylene or a divalent heterocyclic group which has one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, and which optionally forms a fused ring;
20	R <sup>1</sup> , R <sup>2</sup> , R <sup>3</sup> and R <sup>4</sup>	are the same or different and each is a hydrogen atom, a hydroxy, a halogen atom, lower alkoxy, a lower alkyl optionally substituted by a substituent selected from the group consisting of hydroxy, lower alkoxy and halogen atom, or -O-CO-R <sup>11</sup> "
20	-5	wherein R <sup>11</sup> " is lower alkoxy, cycloalkyl, aryl optionally substituted by lower alkyl, or lower alkyl optionally substituted by a substituent selected from the group consisting of acyloxy, aralkyloxycarbonyl and amino optionally substituted by lower alkyl;
	R <sup>5</sup>	is a hydrogen atom, a lower alkyl, or an amino-protecting group; is 1;
25	m R <sup>6</sup>	is an aryl or a cycloalkyl
	••	wherein said aryl and cycloalkyl are optionally substituted by halogen atom or hydroxy; and
30	R <sup>7</sup>	is a hydrogen atom, a lower alkyl optionally substituted by hydroxy or lower alkoxy, an aromatic heterocyclic group which has one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, and which is optionally substituted by lower alkyl, or -CO(Y") <sub>p</sub> R <sup>12</sup> "
		wherein Y" is oxygen atom, sulfur atom or -NR <sup>13</sup> "-wherein R <sup>13</sup> " is hydrogen atom, lower alkyl, hydroxy or amino-protecting group, p is 0 or 1, and R <sup>12</sup> " is hydrogen atom, aralkyl, adamantyl, cycloalkylideneamino, cycloalkyl
35		optionally substituted by lower alkyl, aryl optionally substituted by halogen atom, alkyl optionally substituted by a substitutent selected from the group consisting of hydroxy, lower alkoxy, lower alkoxy, lower alkoxy, lower alkoxy, carboxy, heterocyclic group having one or more hetero atom(s) selected from the group consisting of nitrogen atom, sulfur atom and oxygen atom, and amino optionally substituted by a substituent selected from the group
40		consisting of lower alkyl, aralkyl and amino-protecting group, or heterocyclic group which is optionally substituted by lower alkyl, and which has one or more hetero atom(s) selected from the group consisting of nitrogen atom, sulfur atom and oxygen atom.
45	(4) The amide compound of (1) above, wherein, in the formula (I), at least one symbol selected from the group consisting of R, A, X, M, R <sup>1</sup> , R <sup>2</sup> , R <sup>3</sup> , R <sup>4</sup> , R <sup>5</sup> , m, R <sup>6</sup> and R <sup>7</sup> satisfies the following definitions, and a pharmaceutically acceptable acid addition salt thereof:	
50	R	is a piperazinyl optionally substituted by lower alkyl, a piperidyl optionally substituted by lower alkyl, an amino, or a lower alkoxy substituted by amino wherein amino is optionally substituted by lower alkyl;
-	A X M	is a linear alkylene; is an oxygen atom, a sulfur atom, -NH- or -CH <sub>2</sub> -; is an arylene;
55	R <sup>1</sup> , R <sup>2</sup> , R <sup>3</sup> and R <sup>4</sup>	are the same or different and each is a hydrogen atom, a hydroxy, a halogen atom, or -O-CO-R <sup>11</sup>
- 2		wherein R <sup>11</sup> <sup>m</sup> is lower alkyl optionally substituted by a substituent selected from the group consisting of amino, acyloxy and benzyloxycarbonyl, or phenyl optionally substituted by lower alkyl;

is a hydrogen atom;

 $R^5$ 

	m	is 1;	
	R <sup>6</sup>	is a phenyl; and	
	R <sup>7</sup>	is -COO-R <sup>12</sup> "	
5	wherein R <sup>12</sup> " is hydrogen atom, aralkyl, adamantyl, cyclohexylideneamino, cyclohexyl op		
		ally substituted by lower alkyl, piperidyl optionally substituted by lower alkyl, or alkyl optionally	
		substituted by a substituent selected from the group consisting of hydroxy, lower alkoxy, lower	
	alkoxy lower alkoxy, lower alkoxycarbonyl, acyloxy, piperazinyl and amino optionally subs		
		tuted by lower alkyl.	
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	(5) The amide comp	(5) The amide compound of (4) above, wherein M is phenylene, and a pharmaceutically acceptable acid addition	
	salt thereof.	7	
	(6) The amide comp	ound of (4) above, wherein R <sup>7</sup> is -COO-R <sup>12</sup> wherein R <sup>12</sup> is lower alkyl, or cyclohexyl which	
	is optionally substitu	sted by lower alkyl, and a pharmaceutically acceptable acid addition salt thereof.	
15	(7) The amide compound of (4) above, wherein X is oxygen atom or -CH <sub>2</sub> -, and a pharmaceutically acceptable acid		
	addition salt thereof.		
	(8) The amide compound of (4) above, wherein R <sup>6</sup> is phenyl and m is 1, and a pharmaceutically acceptable acid		
	addition salt thereof	bound of (4) above, wherein R is amino optionally substituted by lower alkyl, piperazinyl option-	
20	ally cubatituted by it	ower alkyl, or piperidyl optionally substituted by lower alkyl, and a pharmaceutically acceptable	
20	acid addition salt the		
	(10) The amide com	spound of (4) above, wherein $R^1$ , $R^2$ , $R^3$ and $R^4$ are the same or different and each is hydrogen	
	atom, hydroxy, halo	ogen atom, or -O-CO-R11" wherein R11" is lower alkyl or phenyl, and a pharmaceutically	
	acceptable acid add		
25		id compound of the formula (I-a)	
		$R^1$ $R^2$	
		$R \longrightarrow A \longrightarrow X \xrightarrow{R^1} \stackrel{R^2}{\longrightarrow} COOH \qquad (I-a)$	
		$R \longrightarrow A \longrightarrow X \longrightarrow M \longrightarrow COOH \qquad (I-a)$	
30		D3 D4	
		$u_*$ $u$	
	wherein;		
35	W. C. C. C.		
	R	is an optionally substituted non-aromatic heterocyclic group containing nitrogen, a hydroxy,	
		$R_a$ , an alkoxy substituted by $R_a$ , an alkylthio substituted by $R_a$ , or an alkylamino substituted by	
		R <sub>a</sub> ,	
		wherein R <sub>a</sub> is amino, guanidino, amidino, carbamoyl, ureido, thioureido, hydrazino, hydrazino	
40		carbonyl or imino, these groups being optionally substituted by a substituent selected from the	
		group consisting of lower alkyl, halogenated lower alkyl, cycloalkyl, aralkyl, aryl and amino-	
		protecting group; is an optionally substituted, linear or branched alkylene which optionally has one or more dou-	
	Α	ble bond(s) or triple bond(s) in the chain, or a single bond;	
45	х	is an oxygen atom, a sulfur atom, a cycloalkylene, a divalent aromatic heterocyclic group hav-	
43	^	ing one ore more hetero atoms selected from the group consisting of a nitrogen atom, sulfur	
		atom and oxygen atom, -SO-, -SO <sub>2</sub> -, -C=C-, -C=C-, -CO-, -COO-, -OOC-, -CS-, -COS-, -O-	
		CO-O-, -NH-CO-NH-, -NH-CS-NH-, -NH-C(=NH)-NH-, -NR8-, -NR8CO-, -CONR8-, -NR8SO <sub>2</sub> -	
		, -SO <sub>2</sub> NR <sup>8</sup> -, -NR <sup>8</sup> -COO-, -OOC-NR <sup>8</sup> -, or -CR <sup>9</sup> R <sup>10</sup> -	
50		wherein R <sup>8</sup> is hydrogen atom, alkyl, cycloalkyl, aryl, aralkyl or amino-protecting group, and R <sup>9</sup>	
		and R <sup>10</sup> are the same or different and each is hydrogen atom, alkyl, cycloalkyl, aryl or aralkyl;	
	M	is an arylene, a cycloalkylene or a divalent heterocyclic group which has one ore more hetero	
		atoms selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom,	
		and which optionally forms a fused ring; and	
55	D1 D2 D3 4 D4	are the same or different and each is a hydrogen atom, a hydroxy, a halogen atom, an alkoxy,	
	$R^1$ , $R^2$ , $R^3$ and $R^4$	are the same of different and sale to a type of the sale	
	K', K', K' and K'	a mercapto, an alkylthio, a nitro, a cyano, a carboxy, an alkoxycarbonyl, an aryloxycarbonyl,	
	R', R', R' and R'	a mercapto, an alkylthio, a nitro, a cyano, a carboxy, an alkoxycarbonyl, an aryloxycarbonyl, an acyl, an alkyl optionally substituted by a substituent selected from the group consisting of	
	K', H-, H- and K	a mercapto, an alkylthio, a nitro, a cyano, a carboxy, an alkoxycarbonyl, an aryloxycarbonyl,	

selected from the group consisting of alkyl, aryl, aralkyl and amino-protecting group, or -O-CO-R<sup>11</sup>

wherein R<sup>11</sup> is optionally substituted alkoxy, optionally substituted aryl, optionally substituted cycloalkyl, optionally substituted aryloxy, optionally substituted aralkyloxy, optionally substituted arylthio, or alkyl optionally substituted by a substitutent selected from the group consisting of alkoxycarbonyl, acyloxy, aryl, aryloxy, aryloxycarbonyl, aralkyloxy, aralkyloxycarbonyl, alkylthio, arylthio, acyl, lower alkoxy, carboxy, halogen atom and amino optionally substituted by lower alkyl or acyl.

(12) The carboxylic acid compound of (11) above, wherein, in the formula (I-a), at least one of R, A, X, M, R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup> and R<sup>4</sup> satisfies the following definitions:

is a piperazinyl optionally substituted by lower alkyl, a piperidyl optionally substituted by lower alkyl, an amino or a lower alkoxy substituted by amino wherein amino is optionally substituted by lower alkyl;

A is a linear alkylene;

X is an oxygen atom, a sulfur atom, -NH- or CH<sub>2</sub>-;

M is an arylene; and

R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup> and R<sup>4</sup> are the same or different and each is a hydrogen atom, a hydroxy, a halogen atom, or -O-CO-R<sup>11</sup>

wherein R<sup>11</sup>" is a lower alkyl optionally substituted by a substituent selected from the group consisting of amino, acyloxy and benzyloxycarbonyl, or phenyl optionally substituted by lower alkyl.

## (13) An amide compound of the formula (I-b)

$$\begin{array}{c|cccc}
R^1 & R^2 & O & (CH_2)_m & R^6 \\
HX & & & & & & \\
R^3 & & & & & & \\
R^4 & & & & & & \\
R^5 & & & & & & \\
\end{array} (I-b)$$

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R

is an oxygen atom, a sulfur atom, a cycloalkylene, a divalent aromatic heterocyclic group having one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, -SO-, -SO<sub>2</sub>-, -C=C-, -C=C-, -CO-, -COO-, -OOC-, -CS-, -COS-, -O-CO-O-, -NH-CO-NH-, -NH-CS-NH-, -NH-C(=NH)-NH-, -NR $^8$ -, -NR $^8$ CO-, -CONR $^8$ -, -NR $^8$ SO<sub>2</sub>-, -SO<sub>2</sub>NR $^8$ -, -NR $^8$ -COO-, -OOC-NR $^8$ - or -CR $^9$ R $^{10}$ -

М

wherein R<sup>8</sup> is hydrogen atom, alkyl, cycloalkyl, aryl, aralkyl or amino-protecting group, and R<sup>9</sup> and R<sup>10</sup> are the same or different and each is hydrogen atom, alkyl, cycloalkyl, aryl or aralkyl; is an arylene, cycloalkylene, or a divalent heterocyclic group which has one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, and which optionally forms a fused ring;

R1, R2, R3 and R4

are the same or different and each is a hydrogen atom, a hydroxy, a halogen atom, an alkoxy, a mercapto, an alkylthio, a nitro, a cyano, a carboxy, an alkoxycarbonyl, an aryloxycarbonyl, an acyl, an alkyl optionally substituted by a substituent selected from the group consisting of hydroxy, lower alkoxy and halogen atom, an amino optionally substituted by a substituent selected from the group consisting of alkyl, aryl, aralkyl and amino-protecting group, or -O-CO-R<sup>11</sup>

wherein R<sup>11</sup> is optionally substituted alkoxy, optionally substituted aryl, optionally substituted cycloalkyl, optionally substituted aryloxy, optionally substituted aralkyloxy, optionally substituted aryloxy, optionally substituted by a substituted arylthio, or alkyl optionally substituted by a substitutent selected from the group consisting of alkoxycarbonyl, acyloxy, aryl, aryloxy, aryloxycarbonyl, aralkyloxy, aralkyloxycarbonyl, alkylthio, arylthio, acyl, lower alkoxy, carboxy, halogen atom and amino optionally substituted by lower alkyl or acyl;

	R <sup>5</sup> is a l	hydrogen atom, an alkyl optionally substituted by halogen atom, optionally substituted
	aralk	yl, or an amino-protecting group;
5	R <sup>6</sup> is an Iowe amin aryl.	or an integer of 1-6; optionally substituted aryl, an optionally substituted cycloalkyl, an optionally substituted ralkyl, an optionally substituted lower alkylthio, an optionally substituted by a substitutent selected from the group consisting of lower alkyl, aralkyl and amino-protecting group, or an optionally substituted heterocyclic group havene or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur
10	atom R <sup>7</sup> is a h subs grou wher	and oxygen atom; and hydrogen atom, an optionally substituted alkyl, an optionally substituted aryl, an optionally stituted aromatic heterocyclic group having one or more hetero atom(s) selected from the p consisting of a nitrogen atom, sulfur atom and oxygen atom, or -CO(Y) <sub>p</sub> R <sup>12</sup> rein Y is oxygen atom, sulfur atom, -NR <sup>13</sup> - or -NR <sup>13</sup> -SO <sub>2</sub> -wherein R <sup>13</sup> is hydrogen atom,
15	alkyl atom cyclo near hydr	, aralkyl, hydroxy, alkoxy, aryl or amino-protecting group, p is 0 or 1, and R is hydrogen in, optionally substituted alkenyl, optionally substituted alkynyl, optionally substituted palkyl, optionally substituted aralkyl, adamantyl, cycloalkylidenino, alkyl optionally substituted by a substituent selected from the group consisting of oxy, alkoxy, alkoxyalkoxy, alkoxycarbonyl, acyloxy, carboxy, heterocyclic group having one
20	or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom oxygen atom, and amino optionally substituted by a substituent selected from the group sisting of alkyl, aryl, aralkyl and amino-protecting group, or optionally substituted heter group having one or more hetero atom(s) selected from the group consisting of a natom, sulfur atom and oxygen atom.	
25	(14) The amide compounconsisting of X, M, R <sup>1</sup> , R <sup>2</sup>	d of (13) above, wherein, in the formula (I-b), at least one symbol selected from the group
	R <sup>3</sup> , R <sup>4</sup> , R <sup>5</sup> , m, R <sup>6</sup> and R	<sup>7</sup> satisfies the following definitions:
	X	is an oxygen atom, a sulfur atom or -NH-;
30	M	is an arylene;
	$R^1$ , $R^2$ , $R^3$ and $R^4$	are the same or different and each is a hydrogen atom, a hydroxy, a halogen atom, or -
		O-CO-R <sup>11</sup> " wherein R <sup>11</sup> " is lower alkyl optionally substituted by a substituent selected from the group consisting of amino, acyloxy and benzyloxycarbonyl, or a phenyl optionally substi-
35	R <sup>5</sup>	tuted by lower alkyl;
	m	is a hydrogen atom; is 1;
	. R <sup>6</sup>	is a phenyl; and
	R <sup>7</sup>	is -COO-R <sup>12</sup> **
		wherein R <sup>12</sup> " is hydrogen atom, aralkyl, adamantyl, cyclohexylideneamino, piperidyl optionally substituted by lower alkyl, cyclohexyl optionally substituted by lower alkyl, or alkyl optionally substituted by a substituent selected from the group consisting of hydroxy, lower alkoxy, lower alkoxy, lower alkoxy, lower alkoxy, acyloxy, piperazinyl, and amino optionally substituted by lower alkyl.
45	any one of (1) to (10) about (16) An inflammatory cy	omposition comprising a pharmaceutically acceptable carrier, and the amide compound of ove or a pharmaceutically acceptable acid addition salt thereof. tokine production suppressor comprising the amide compound of any one of (1) to (10)
	above or a pharmaceutic	ally acceptable acid addition salt thereof as an active ingredient.
50	(17) An agent for the trea	tment or prophylaxis of an inflammatory diseases, comprising the amide compound of any

In the present specification, each substituent means as follows.

"Alkoxy" is linear or branched alkoxy having 1 to 6 carbon atoms, which is exemplified by methoxy, ethoxy, propoxy, sopropoxy, butoxy, isobutoxy, sec-butoxy, tert-butoxy, pentyloxy, isopentyloxy, neopentyloxy, tert-pentyloxy, hexyloxy, isohexyloxy and neohexyloxy, with preference given to linear or branched alkoxy having 1 to 4 carbon atoms, such as methoxy, ethoxy, propoxy, isopropoxy, butoxy, isobutoxy, sec-butoxy and tert-butoxy.

one of (1) to (10) above or a pharmaceutically acceptable acid addition salt thereof as an active ingredient.

"Lower alkoxy" is linear or branched alkoxy having 1 to 4 carbon atoms, which is exemplified by methoxy, ethoxy,

propoxy, isopropoxy, butoxy, isobutoxy, sec-butoxy and tert-butoxy, with preference given to methoxy and ethoxy.

"Alkytthio" is linear or branched alkytthio having 1 to 6 carbon atoms, which is exemplified by methytthio, ethytthio, propytthio, isopropytthio, butytthio, isobutythio, sec-butythio, tert-butythio, pentythio, isopentythio, neopentythio, tert-pentythio, hexytthio, isohexytthio and neohexytthio.

"Lower alkytthio" is linear or branched alkytthio having 1 to 4 carbon atoms, which is exemplified by methytthio, ethylthio, propylthio, isopropylthio, butylthio, isobutylthio, sec-butylthio and tert-butylthio.

"Alkylamino" is linear or branched, monoalkylamino or dialkylamino which has 1 to 6 carbon atoms, which is exemplified by methylamino, dimethylamino, ethylamino, diethylamino, methylethylamino, propylamino, isopropylamino, butylamino, isobutylamino, sec-butylamino, tert-butylamino, pentylamino, isopentylamino, neopentylamino, tert-pentylamino, hexylamino, isohexylamino and neohexylamino, with preference given to linear alkylamino, such as methylamino, dimethylamino, ethylamino, propylamino, butylamino, pentylamino and hexylamino. Particularly preferred is linear alkylamino having 1 to 4 carbon atoms, which is exemplified by methylamino, dimethylamino, ethylamino, diethylamino, propylamino and butylamino.

"Non-aromatic heterocyclic group containing nitrogen" is 3- to 7-membered non-aromatic heterocyclic group which has at least one nitrogen atom and optionally a sulfur atom or oxygen atom, and which is optionally fused with benzene ring. Specific examples thereof include aziridinyl, thiazetidinyl, azetidinyl, pyrrolidinyl, pyrrolinyl, imidazolinyl, pyrazolidinyl, pyrazolidinyl, morpholiny, morpholino, oxazinyl, thiazinyl, piperazinyl, piperidyl, piperidyl, piperidyl, azepinyl, thiazepinyl, diazepinyl, perhydrodiazepinyl, azepinyl, perhydroazepinyl, indolinyl and isoindolinyl. Preferred are aziridinyl, azetidinyl, pyrrolidinyl, pyrazolidinyl, morpholinyl, morpholino, piperazinyl, piperidyl, piperidino and perhydroazepinyl, and particularly preferred are pyrrolidinyl, morpholino, piperazinyl, piperidyl and piperidino.

"Alkyl" is linear or branched alkyl having 1 to 6 carbon atoms, which is exemplified by methyl, ethyl, propyl, isopropyl, butyl, isobutyl, sec-butyl, tert-butyl, pentyl, isopentyl, neopentyl, tert-pentyl, hexyl, isohexyl and neohexyl.

"Lower alkyl" is linear or branched alkyl having 1 to 4 carbon atoms, which is exemplified by methyl, ethyl, propyl, isopropyl, butyl, isobutyl, sec-butyl and tert-butyl.

"Halogen atom" is specifically a fluorine atom, chlorine atom, bromine atom or iodine atom.

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"Halogenated lower alkyl" is that wherein the above-mentioned lower alkyl is substituted by a halogen atom, and is exemplified by fluoromethyl, chloromethyl, bromomethyl, difluoromethyl, dichloromethyl, trifluoromethyl, trichloromethyl, difluoroethyl, dichloroethyl, pentatrifluoroethyl, trichloroethyl and fluoropropyl, with preference given to fluoromethyl, chloromethyl, difluoromethyl, dichloromethyl and trifluoromethyl.

"Cycloalkyl" is that having 3 to 7 carbon atoms, which is exemplified by cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl and cycloheptyl, with preference given to cycloalkyl having 5 or 6 carbon atoms, such as cyclopentyl and cyclohexyl.

"Aralkyl" is that wherein alkyl is substituted by aryl and is exemplified by benzyl, benzhydryl, trityl, phenethyl, 3-phenylpropyl, 2-phenylpropyl, 4-phenylbutyl and naphthylmethyl, with preference given to benzyl and phenethyl.

"Aralkyloxy" is that having the above-mentioned aralkyl, which is exemplified by benzyloxy, benzhydryloxy, trityloxy, phenethyloxy, 3-phenylpropyloxy, 2-phenylpropyloxy, 4-phenylbutyloxy and naphthylmethoxy, with preference given to benzyloxy and phenethyloxy.

"Aralkyloxycarbonyl" is that having the above-mentioned aralkyl, which is exemplified by benzyloxycarbonyl, benzhydryloxycarbonyl, trityloxycarbonyl, phenethyloxycarbonyl, 3-phenylpropyloxycarbonyl, 2-phenylpropyloxycarbonyl, 4-phenylbutyloxycarbonyl and naphthylmethoxycarbonyl, with preference given to benzyloxycarbonyl and phenethyloxycarbonyl.

"Aryl" is phenyl, naphthyl, anthryl, phenanthryl or biphenyl, with preference given to phenyl and naphthyl.

"Aryloxy" is that having the above-mentioned aryl, which is exemplified by phenoxy and naphthyloxy.

"Aryloxycarbonyl" is that having the above-mentioned aryl, which is exemplified by phenoxycarbonyl and naphthyloxycarbonyl.

"Arylthio" is that having the above-mentioned aryl, which is exemplified by phenylthio and naphthylthio.

"Amino-protecting group" is a protecting group conventionally used, which is subject to no particular limitation as long as it protects amino from various reactions. Specific examples include acyl such as formyl, acetyl, propionyl, butyryl, oxalyl, succinyl, pivaloyl, 2-chloroacetyl, 2-bromoacetyl, 2-iodoacetyl, 2,2-dichloroacetyl, 2,2,2-trichloroacetyl, 2,2,2-trifluoroacetyl, phenoxyacetyl, benzoyl, 4-chlorobenzoyl, 4-methoxybenzoyl, 4-nitrobenzoyl, naphthylcarbonyl, adamantylcarbonyl and phthaloyl; alkoxycarbonyl such as methoxycarbonyl, ethoxycarbonyl, propoxycarbonyl, isopropoxycarbonyl, tert-butoxycarbonyl, pentyloxycarbonyl, isopentyloxycarbonyl, cyclohexyloxycarbonyl, 2-chloroethoxycarbonyl, 2-iodoethoxycarbonyl, 2,2,2-trichloroethoxycarbonyl, 2,2,2-trichloro-tert-butoxycarbonyl, benzhydryloxycarbonyl, 2-iodoethoxycarbonyl, pentyloxycarbonyl, phenacyloxycarbonyl, 2-trimethylsilylethoxycarbonyl, 2-triphenylsilylethoxycarbonyl and fluorenyl-9-methoxycarbonyl; alkenyloxycarbonyl such as vinyloxycarbonyl, 2-propenyloxycarbonyl, 2-chloro-2-propenyloxycarbonyl, 3-methoxycarbonyl; aralkyloxycarbonyl, such as benzyloxycarbonyl, 4-bromobenzyloxycarbonyl, 2-chlorobenzyloxycarbonyl, 3-chlorobenzyloxycarbonyl, 3,5-dimethoxybenzyloxycarbonyl, 4-bromobenzyloxycarbonyl, 2-chlorobenzyloxycarbonyl, 3-chlorobenzyloxycarbonyl, 3,5-dimethoxybenzyloxycarbonyl, 4-bromobenzyloxycarbonyl, 2-chlorobenzyloxycarbonyl, 3-chlorobenzyloxycarbonyl, 3,5-dimethoxybenzyloxycarbonyl, 4-bromobenzyloxycarbonyl, 4-bromobenzyloxycarbonyl, 4-bromobenzyloxycarbonyl, 4-bromobenzyloxycarbonyl, 4-bromobenzyloxycarbonyl, 4-bromobenzyloxycarbonyl, 4-bromobenzyloxycarbonyl, 4-bromobenzyloxycarbonyl, 4-bromobenzyloxyc

methoxybenzyloxycarbonyl, 2-nitrobenzyloxycarbonyl, 4-nitrobenzyloxycarbonyl, 2-nitro-4,5-dimethoxybenzyloxycarbonyl, 3,4,5-trimethoxybenzyloxycarbonyl and phenethyloxycarbonyl; lower alkylsilyl such as trimethylsilyl and tert-butyldimethylsilyl; alkylenebis(dialkylsilyl) such as ethylenebis(dimethylsilyl), propylenebis(dimethylsilyl) and ethylenebis(diethylsilyl); alkylthiocarbonyl such as methylthiocarbonyl, ethylthiocarbonyl, butylthiocarbonyl and tert-butylthiocarbonyl; aralkylthiocarbonyl such as benzylthiocarbonyl; phosphoryl such as dicyclohexylphosphoryl, diphenylphosphoryl, dibenzylphosphoryl, di-(4-nitrobenzyl)phosphoryl and phenoxyphenylphosphoryl; and phosphinyl such as diethylphosphinyl, diphenylphosphinyl.

"Linear or branched alkylene optionally having one or more double bond(s) or triple bond(s) in the chain" is linear or branched alkylene having 1 to 10 carbon atoms, which may have one ore more double bonds or triple bonds in the chain, and is exemplified by methylene, ethylene, trimethylene, tetramethylene, pentamethylene, hexamethylene, heptamethylene, octamethylene, nonamethylene, decamethylene, dimethylmethylene, diethylmethylene, propylene, methylethylene, ethylethylene, propylethylene, isopropylethylene, methylpentaethylene, ethylethylene, propylethylene, isopropylethylene, methylpentaethylene, ethylethylene, dimethyltrimethylene, vinylene, propenylene, butenylene, butadienylene, pentenylene, pentadienylene, hexadienylene, hexadienylene, hexatrienylene, heptadienylene, heptatrienylene, octaylene, octadienylene, octateraenylene, propynylene, butynylene, pentynylene and methylpropynylene, with preference given to linear alkylene, such as methylene, ethylene, trimethylene, tetramethylene, pentamethylene, hexamethylene, hexamethylene, propynylene, butadienylene, pentenylene, octamethylene, nonamethylene, decamethylene, vinylene, propenylene, butenylene, butadienylene, pentenylene, pentadienylene, hexamethylene, hexamethylene, propynylene, butynylene and pentynylene. Particularly preferred is linear alkylene having 1 to 8 carbon atoms, such as methylene, ethylene, trimethylene, tetramethylene, pentamethylene, hexamethylene, heptamethylene.

"Divalent aromatic heterocyclic group having one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom" is 5- or 6-membered divalent aromatic heterocyclic group having one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, which is exemplified by divalent groups of tetrazole ring, oxadiazole ring, thiadiazole ring, triazole ring, oxazole ring, isothiazole ring, imidazole ring, pyrazole ring, pyrrole ring, furan ring, thiophene ring, tetrazine ring, triazine ring, pyrazine ring, pyridazine ring, pyrimidine ring and pyridine ring. Preferred is 5-membered divalent aromatic heterocyclic group, which is exemplified by divalent groups of tetrazole ring, oxadiazole ring, thiadiazole ring, triazole ring, oxazole ring, isoxazole ring, isothiazole ring, imidazole ring, pyrrole ring, pyrrole ring, furan ring and thiophene ring. Particularly preferred are divalent groups of oxadiazole ring, thiadiazole ring and triazole ring.

"Cycloalkylene" is that having 3 to 7 carbon atoms, namely, divalent cycloalkyl, which is specifically exemplified by cyclopropylene, cyclobutylene, cyclopentylene, cyclohexylene and cycloheptylene. Preferred is cycloalkylene having 5 or 6 carbon atoms, which is exemplified by cyclopentylene and cyclohexylene.

"Arylene" is exemplified by phenylene, naphthylene, anthrylene, phenanthrylene and biphenylene, with preference given to phenylene, naphthylene and biphenylene.

"Divalent heterocyclic group which has one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, and which optionally forms a fused ring" is specifically exemplified by divalent heterocyclic groups of dioxolane ring, dithiol ring, pyrrolidine ring, morpholine ring, oxazine ring, piperazine ring, piperidine ring, pyrroline ring, imidazolidine ring, imidazoline ring, pyrazolidine ring, pyrazoline ring, thiatriazole ring, tetrazole ring, oxadiazole ring, thiadiazole ring, triazole ring, isoxazole ring, oxazole ring, thiazole ring, imidazole ring, pyrazole ring, pyrrole ring, furan ring, thiophene ring, tetrazine ring, triazine ring, pyrazine ring, pyridazine ring, pyr ine ring, furoisoxazole ring, imidazothiazole ring, thienoisothiazole ring, thienothiazole ring, imidazopyrazole ring, cyclopentapyrazole ring, pyrrolopyrrole ring, thienothiophene ring, thiadiazolopyrimidine ring, thiazolothiazine ring, thiazolopyrimidine ring, thiazolopyridine ring, oxazolopyrimidine ring, oxazolopyridine ring, benzoxazole ring, benzisothiazole ring, benzothiazole ring, imidazopyrazine ring, purine ring, pyrazolopyrimidine ring, imidazopyridine ring, benzimidazole ring, indazole ring, benzoxathiole ring, benzodioxole ring, benzodithiol ring, indolizine ring, indoline ring, isoindoline ring, furopyrimidine ring, furopyridine ring, benzofuran ring, isobenzofuran ring, thienopyrimidine ring, thienopyridine ring, benzothiophene ring, cyclopentaoxazine ring, cyclopentafuran ring, benzoxazine ring, benzothiazine ring, quinazoline ring, naphthyridine ring, quinoline ring, isoquinoline ring, benzopyran ring, pyridopyridazine ring and pyridopyrimidine ring. Preferred are divalent heterocyclic groups of piperazine ring, piperidine ring, pyridine ring, benzoxazole ring, benzisothiazole ring, benzothiazole ring and benzimidazole ring.

"Alkoxycarbonyl" is linear or branched alkoxycarbonyl having 2 to 7 carbon atoms, which is exemplified by methoxycarbonyl, ethoxycarbonyl, propoxycarbonyl, isopropoxycarbonyl, butoxycarbonyl, isobutoxycarbonyl, sec-butoxycarbonyl, tert-butoxycarbonyl, pentyloxycarbonyl, isopentyloxycarbonyl, neopentyloxycarbonyl, tert-pentyloxycarbonyl, hexyloxycarbonyl, isohexyloxycarbonyl and neohexyloxycarbonyl, with preference given to linear or branched alkoxycarbonyl having 2 to 5 carbon atoms, such as methoxycarbonyl, ethoxycarbonyl, propoxycarbonyl, isopropoxycarbonyl, butoxycarbonyl, isobutoxycarbonyl, sec-butoxycarbonyl and tert-butoxycarbonyl.

"Lower alkoxycarbonyl" is linear or branched alkoxycarbonyl having 2 to 5 carbon atoms, which is exemplified by

methoxycarbonyl, ethoxycarbonyl, propoxycarbonyl, isopropoxycarbonyl, butoxycarbonyl, isobutoxycarbonyl, secbutoxycarbonyl and tert-butoxycarbonyl, with preference given to methoxycarbonyl and ethoxycarbonyl.

"Acyl" specifically means, for example, formyl, acetyl, propionyl, butyryl, isobutyryl, valeryl, isovaleryl, pivaloyl, caproyl, isocaproyl, acryloyl, propioloyl, methacryloyl, crotonoyl, isocrotonoyl, benzoyl, naphthoyl, toluoyl, hydroatropoyl, atropoyl, cinnamoyl, furoyl, glyceroyl, tropoyl, benziloyl, salicyloyl, anisoyl, vanilloyl, veratroyl, piperonyloyl, protocatechuoyl or galloyl, with preference given to formyl, acetyl, propionyl, butyryl, isobutyryl, valeryl, isovaleryl, pivaloyl, benzoyl and naphthoyl.

"Acyloxy" is that having the above-mentioned acyl, which is exemplified by formyloxy, acetyloxy, propionyloxy, butyryloxy, isobutyryloxy, valeryloxy, isovaleryloxy, pivaloyloxy, caproyloxy, isocaproyloxy, acryloyloxy, propioloyloxy, methacryloyloxy, crotonoyloxy, isocrotonoyloxy, benzoyloxy, naphthoyloxy, toluoyloxy, hydroatropoyloxy, atropoyloxy, cinnamoyloxy, furoyloxy, glyceroyloxy, tropoyloxy, benziloyloxy, salicyloyloxy, anisoyloxy, vanilloyloxy, veratroyloxy, piperonyloyloxy, protocatechuoyloxy and galloyloxy, with preference given to formyloxy, acetyloxy, propionyloxy, butyryloxy, isobutyryloxy, valeryloxy, isovaleryloxy, pivaloyloxy, benzoyloxy and naphthoyloxy.

"Heterocyclic group having one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom" at R<sup>6</sup> is 3- to 7-membered heterocyclic group having one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, which is exemplified by aziridinyl, oxiranyl, azetyl, azetidinyl, oxetanyl, thiatriazolyl, tetrazolyl, dithiazolyl, oxadiazolyl, thiadiazolyl, triazolyl, oxazolyl, isooxazolyl, thiazolyl, isothiazolyl, imidazolyl, pyrazolyl, dioxolanyl, pyrrolyl, pyrrolidinyl, furanyl, thienyl, tetrazinyl, dithiadiazinyl, thiadiazinyl, triazinyl, morpholinyl, morpholino, oxazinyl, thiazinyl, piperazinyl, pyrazinyl, pyridazinyl, pyrimidinyl, piperidyl, piperidino, pyridyl, pyranyl, thiopyranyl, dioxazepinyl, diazepinyl and azepinyl. Preferred is 5- or 6-membered heterocyclic group, which is exemplified by thiatriazolyl, tetrazolyl, dithiazolyl, oxadiazolyl, triazolyl, triazolyl, oxazolyl, isooxazolyl, thiazolyl, isothiazolyl, imidazolyl, pyrazolyl, dioxolanyl, pyrrolyl, pyrrolidinyl, furanyl, thienyl, tetrazinyl, dithiadiazinyl, thiadiazinyl, triazinyl, morpholinyl, morpholino, oxazinyl, thiazinyl, piperazinyl, pyrazinyl, pyrazinyl, pyrazinyl, pyrazinyl, piperidino, pyridyl, pyranyl and thiopyranyl. Particularly preferred are pyrrolyl, furanyl, thienyl, piperazinyl, piperidino and pyridyl.

"Aromatic heterocyclic group having one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen" is 5- or 6-membered aromatic heterocyclic group having one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, which is exemplified by tetrazolyl, oxadiazolyl, thiadiazolyl, triazolyl, oxazolyl, isooxazolyl, thiazolyl, isothiazolyl, imidazolyl, pyrazolyl, pyrrolyl, furanyl, thienyl, tetrazinyl, triazinyl, pyrazinyl, pyridazinyl, pyridiazinyl, pyridiazinyl, pyridiazinyl, thiadiazolyl, triazolyl, oxazolyl, isooxazolyl, thiadiazolyl, triazolyl, isooxazolyl, thiadiazolyl, isothiazolyl, imidazolyl, pyrrolyl, furanyl and thienyl. Particularly preferred are oxadiazolyl, thiadiazolyl and triazolyl.

"Alkoxyalkoxy" is that wherein linear or branched alkoxy having 1 to 6 carbon atoms has been substituted by linear or branched alkoxy having 1 to 6 carbon atoms, and is exemplified by methoxymethoxy, ethoxymethoxy, propoxymethoxy, isopropoxymethoxy, butoxymethoxy, isobutoxymethoxy, sec-butoxymethoxy, tert-butoxymethoxy, pentyloxymethoxy, isopentyloxymethoxy, neopentyloxymethoxy, tert-pentyloxymethoxy, hexyloxymethoxy, isohexyloxymethoxy, neohexyloxymethoxy, tert-hexyloxymethoxy, methoxyethoxy, ethoxyethoxy, propoxyethoxy, isopropoxyethoxy, butoxyethoxy, isobutoxyethoxy, sec-butoxyethoxy, tert-butoxyethoxy, pentyloxyethoxy, isopentyloxyethoxy, neopentyloxyethoxy, tert-pentyloxyethoxy, hexyloxyethoxy, isohexyloxyethoxy, neohexyloxyethoxy, tert-hexyloxyethoxy, methoxypropoxy, ethoxypropoxy, propoxypropoxy, isopropoxypropoxy, butoxypropoxy, isobutoxypropoxy, sec-butoxypropoxy, tert-butaxypropoxy, pentyloxypropoxy, isopentyloxypropoxy, neopentyloxypropoxy, tert-pentyloxypropoxy, hexyloxypropoxy, isohexyloxypropoxy, neohexyloxypropoxy, tert-hexyloxypropoxy, methoxybutoxy, ethoxybutoxy, propoxybutoxy, isopropoxybutoxy, butoxybutoxy, isobutoxybutoxy, sec-butoxybutoxy, tert-butoxybutoxy, pentyloxybutoxy, isopentyloxybutoxy, neopentyloxybutoxy, tert-pentyloxybutoxy, hexyloxybutoxy, isohexyloxybutoxy, neohexyloxybutoxy, tert-hexyloxybutoxy, methoxypentyloxy, ethoxypentyloxy, propoxypentyloxy, isopropoxypentyloxy, butoxypentyloxy, isobutoxypentyloxy, sec-butoxypentyloxy, tert-butoxypentyloxy, pentyloxypentyloxy, isopentyloxypentyloxy, neopentyloxypentyloxy, tert-pentyloxypentyloxy, hexyloxypentyloxy, isohexyloxypentyloxy, neohexyloxypentyloxy, terthexyloxypentyloxy, methoxyhexyloxy, ethoxyhexyloxy, propoxyhexyloxy, isopropoxyhexyloxy, butoxyhexyloxy, isobutoxyhexyloxy, sec-butoxyhexyloxy, tert-butoxyhexyloxy, pentyloxyhexyloxy, isopentyloxyhexyloxy, neopentyloxyhexyloxy, tert-pentyloxyhexyloxy, hexyloxyhexyloxy, isohexyloxyhexyloxy, neohexyloxyhexyloxy and tert-hexyloxyhexyloxy. Preferred is that wherein linear or branched alkoxy having 1 to 4 carbon atoms has been substituted by linear or branched alkoxy having 1 to 4 carbon atoms, and is exemplified by methoxymethoxy, ethoxymethoxy, propoxymethoxy, isopropoxymethoxy, butoxymethoxy, isobutoxymethoxy, sec-butoxymethoxy, tert-butoxymethoxy, methoxyethoxy, ethoxyethoxy, propoxyethoxy, isopropoxyethoxy, butoxyethoxy, isobutoxyethoxy, sec-butoxyethoxy, tert-butoxyethoxy, methoxypropoxy, ethoxypropoxy, propoxypropoxy, isopropoxypropoxy, butoxypropoxy, isobutoxypropoxy, sec-butoxypropoxy, tert-butoxypropoxy, methoxybutoxy, ethoxybutoxy, propoxybutoxy, isopropoxybutoxy, butoxybutoxy, isobutoxybutoxy, sec-butoxybutoxy and tert-butoxybutoxy.

"Lower alkoxy lower alkoxy" is that wherein linear or branched alkoxy having 1 to 4 carbon atoms has been substi-

tuted by linear or branched alkoxy having 1 to 4 carbon atoms, and is exemplified by methoxymethoxy, ethoxymethoxy, propoxymethoxy, butoxymethoxy, isobutoxymethoxy, sec-butoxymethoxy, tert-butoxymethoxy, methoxyethoxy, propoxyethoxy, isobutoxyethoxy, isobutoxyethoxy, sec-butoxyethoxy, tert-butoxyethoxy, propoxyethoxy, propoxypropoxy, isopropoxypropoxy, butoxypropoxy, isobutoxypropoxy, sec-butoxypropoxy, methoxybutoxy, ethoxybutoxy, propoxybutoxy, isopropoxybutoxy, butoxybutoxy, isobutoxybutoxy, butoxybutoxy, isobutoxybutoxy, ethoxybutoxy, with preference given to methoxymethoxy, ethoxymethoxy, methoxyethoxy and ethoxyethoxy.

"Heterocyclic group having one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen" at R<sup>12</sup> means 3- to 7-membered heterocyclic group having one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen, and is exemplified by aziridinyl, oxiranyl, azetyl, azetidinyl, oxetanyl, thiatriazolyl, tetrazolyl, dithiazolyl, oxadiazolyl, thiadiazolyl, triazolyl, isooxazolyl, thiazolyl, isothiazolyl, imidazolyl, pyrazolyl, dioxolanyl, pyrrolyl, pyrrolidinyl, furanyl, thienyl, tetrazinyl, dithiadiazinyl, triazinyl, morpholinyl, morpholino, oxazinyl, thiazinyl, piperazinyl, pyrazinyl, pyrimidinyl, piperidino, pyridyl, pyranyl, thiopyranyl, dioxazepinyl, diazepinyl and azepinyl. Preferred is 5- or 6-membered heterocyclic group, such as thiatriazolyl, tetrazolyl, dithiazolyl, oxadiazolyl, thiadiazolyl, triazolyl, isooxazolyl, thiazolyl, isoothiazolyl, imidazolyl, pyrazolyl, dioxolanyl, pyrrolyl, pyrrolidinyl, furanyl, thienyl, tetrazinyl, dithiadiazinyl, thiadiazinyl, triazinyl, morpholinyl, morpholino, oxazinyl, thiazinyl, piperazinyl, pyrazinyl, pyrimidinyl, piperidyl, piperidyl,

"Alkenyl" is linear or branched alkenyl having 2 to 6 carbon atoms, which is exemplified by allyl, vinyl, propenyl, iso-propenyl, 1-methyl-2-propenyl, 2-methyl-2-propenyl, 1-methyl-1-butenyl, crotyl, 1-methyl-3-butenyl, 3-methyl-2-butenyl, 1-pentenyl, 1-methyl-2-pentenyl, 4-pentenyl, 1-hexenyl, 3-hexenyl and 4-hexenyl.

"Alkynyl" is linear or branched alkynyl having 2 to 6 carbon atoms, which is exemplified by propargyl, 2-butynyl, 1-methyl-2-butynyl, 2-pentynyl, 1-methyl-3-pentynyl, 1-methyl-4-pentynyl, 1-hexynyl and 5-hexynyl.

"Cycloalkylideneamino" specifically means cyclopropylideneamino, cyclobutylideneamino, cyclopentylideneamino, cyclopentylideneamino, cyclopentylideneamino and cyclohexylideneamino, with preference given to cyclopentylideneamino and cyclohexylideneamino.

"Alkoxy" of the substituted alkoxy at R is linear or branched alkoxy having 1 to 6 carbon atoms, which is exemplified by methoxy, ethoxy, propoxy, isopropoxy, butoxy, isobutoxy, sec-butoxy, tert-butoxy, pentyloxy, isopentyloxy, neopentyloxy, tert-pentyloxy, hexyloxy, isohexyloxy and neohexyloxy, with preference given to linear alkoxy, such as methoxy, ethoxy, propoxy, butoxy, pentyloxy and hexyloxy. Particularly preferred is linear alkoxy having 1 to 4 carbon atoms, which is exemplified by methoxy, ethoxy, propoxy and butoxy.

"Alkylthio" of the substituted alkylthio at R is linear or branched alkylthio having 1 to 6 carbon atoms, which is exemplified by methylthio, ethylthio, propylthio, isopropylthio, butylthio, isobutylthio, sec-butylthio, tert-butylthio, pentylthio, isopentylthio, neopentylthio, tert-pentylthio, hexylthio, isohexylthio and neohexylthio, with preference given to linear alkylthio such as methylthio, ethylthio, propylthio, butylthio, pentylthio and hexylthio. Particularly preferred is linear alkylthio having 1 to 4 carbon atoms, which is exemplified by methylthio, ethylthio, propylthio and butylthio.

"Optionally substituted" of "optionally substituted non-aromatic heterocyclic group containing nitrogen" means that the group may be substituted by 1 to 3 substituent(s) and said substituents may be the same or different. The position of the substituent(s) is optional and is not particularly limited. Specific examples of the substituents include the above-mentioned lower alkyl, the above-mentioned halogenated lower alkyl, the above-mentioned cycloalkyl, the above-mentioned aralkyl, the above-mentioned aryl, and the above-mentioned amino-protecting group. Preferred are lower alkyl and amino-protecting group.

"Optionally substituted" of "optionally substituted linear or branched alkylene which may have one or more double bond(s) or triple bond(s) in the chain" means that the group may be substituted by one or more substituent(s). Examples of the substituents include the above-mentioned halogen atom, hydroxy, amino which may be substituted by a substituent selected from the group consisting of the above-mentioned lower alkyl, the above-mentioned halogenated lower alkyl, the above-mentioned cycloalkyl, the above-mentioned aralkyl, the above-mentioned aryl and the above-mentioned amino protecting group, the above-mentioned lower alkoxy, the above-mentioned aralkyl and the above-mentioned cycloalkyl.

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"Optionally substituted" of "optionally substituted alkoxy" and "optionally substituted alkylthio" at R<sup>11</sup> means that the group may be substituted by one or more substituent(s), and said substituents may be the same or different. The position of the substituent(s) is optional and is not particularly limited. Specific examples of the substituents include the above-mentioned halogen atom, the above-mentioned lower alkoxy, the above-mentioned alkylthio, amino which may be substituted by the above-mentioned lower alkyl or the above-mentioned acyl, carboxy, the above-mentioned alkoxy-carbonyl, the above-mentioned acyl, the above-mentioned acyl, the above-mentioned aryloxy, the above-mentioned aralkyloxy, and the above-mentioned aralkyloxycarbonyl. Preferred are amino, lower alkoxy, halogen atom, carboxy, alkoxycarbonyl and aralkyloxycarbonyl.

"Optionally substituted" of "optionally substituted aryl", "optionally substituted cycloalkyl", "optionally substituted aryloxy", "optionally substituted arylthio" at R<sup>11</sup> means that they may have 1 t 3 substituent(s) on the ring wherein said substituents may be the same or different. The position of the substituent(s) is optional and is not particularly limited. Specific examples of th substituents include the above-mentioned lower alkyl, the above-mentioned halogen atom, the above-mentioned lower alkoxy, the above-mentioned alkylthio, amino which may be substituted by the above-mentioned lower alkyl or the above-mentioned acyl, carboxy, the above-mentioned alkoxycarbonyl, the above-mentioned acyl, the above-mentioned arylthio, the above-mentioned aryloxy, the above-mentioned arylthio, the above-mentioned aryloxycarbonyl, the above-mentioned aralkyloxycarbonyl. Preferred are lower alkyl, amino, lower alkoxy, halogen atom, carboxy, alkoxycarbonyl and aralkyloxycarbonyl. Particularly preferred is lower alkyl.

"Optionally substituted" of "optionally substituted aralky!" at R<sup>5</sup> means that it may have 1 to 3 substituent(s) on aryl wherein said substituents may be the same or different. The position of the substituent(s) is optional and is not particularly limited. Specific examples of the substituents include the above-mentioned lower alkyl, the above-mentioned lower alkyl or the above-mentioned acyl, amino which may be substituted by the above-mentioned lower alkyl or the above-mentioned acyl, the above-mentioned alkoxycarbonyl, the above-mentioned aryloxy, the above-mentioned alkylthio, the above-mentioned aryloxy, the above-mentioned aryloxy. Preferred are lower alkyl, lower alkoxy and halogen atom. Preferred is lower alkyl.

"Optionally substituted" of "optionally substituted lower alkyl", "optionally substituted lower alkoxy" and "optionally substituted lower alkylthio" at R<sup>6</sup> means that the group may be substituted by one or more substituent(s) and said substituents may be the same or different. The position of the substituent(s) is optional and is not particularly limited. Specific examples of the substituents include the above-mentioned halogen atom, hydroxy, the above-mentioned alkoxy, the above-mentioned aryloxy, amino which may be substituted by the above-mentioned lower alkyl or the above-mentioned acyl, mercapto, the above-mentioned alkylthio, the above-mentioned arylthio, carboxy, the above-mentioned alkoxycarbonyl, the above-mentioned halogenated lower alkyl, sulfamoyl, cyano, nitro, alkylsulfonyl such as methylsulfonyl, ethylsulfonyl, isopropylsulfonyl, alkylsulfinyl such as methylsulfonyl, ethylsulfonyl, ethylsulfonyl, Preferred are halogen atom, hydroxy, alkoxy, amino, carboxy and alkoxycarbonyl.

"Optionally substituted" of "optionally substituted aryl", "optionally substituted cycloalkyl" and "optionally substituted heterocyclic group having one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom" at R<sup>6</sup> means that the group may be substituted by one or more substituent(s) and said substituents may be the same or different. The position of the substituent(s) is optional and is not particularly limited. Specific examples of the substituents include the above-mentioned lower alkyl, the above-mentioned halogen atom, hydroxy, the above-mentioned alkoxy, the above-mentioned aryloxy, amino which may be substituted by the above-mentioned lower alkyl or the above-mentioned acyl, mercapto, the above-mentioned alkylthio, the above-mentioned aryloxycarbonyl, carbamoyl, the above-mentioned halogenated lower alkyl, sulfamoyl, cyano, nitro, alkylsulfonyl such as methylsulfonyl, ethylsulfonyl and isopropylsulfonyl, alkylsulfinyl such as methylsulfonyl such as phenylsulfonyl. Preferred are lower alkyl, halogen atom, hydroxy, alkoxy, amino, carboxy and alkoxycarbonyl.

"Optionally substituted" of "optionally substituted alkyl" at R<sup>7</sup> means that the group may be substituted by one or more substituent(s) and said substituent(s) may be the same or different. The position of the substituent(s) is optional and is not particularly limited. Specific examples of the substituents include hydroxy, the above-mentioned lower alkoxy, mercapto, the above-mentioned lower alkylthio, carboxy, the above-mentioned lower alkoxycarbonyl, halogen atom, and amino which may be substituted by the above-mentioned lower alkyl or the above-mentioned acyl. Preferred are hydroxy, halogen atom and lower alkoxy.

"Optionally substituted" of "optionally substituted aryl" and "optionally substituted aromatic heterocyclic group having one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom" at R<sup>7</sup> means that they may have 1 to 3 substituent(s) on the ring wherein said substituents may be the same or different. The position of the substituent(s) is optional and is not particularly limited. Specific examples of the substituents include the above-mentioned lower alkyl, hydroxy, the above-mentioned lower alkoxy, mercapto, the above-mentioned lower alkylthio, carboxy, the above-mentioned lower alkoxycarbonyl, halogen atom, and amino which may be substituted by the above-mentioned lower alkyl or the above-mentioned acyl. Preferred are hydroxy, lower alkyl, halogen atom and lower alkoxy.

"Optionally substituted" of "optionally substituted alkenyl" and "optionally substituted alkynyl" at R<sup>12</sup> means that the group may be substituted by one or more substituent(s) and said substituent(s) may be the same or different. The position of the substituent(s) is optional and is not particularly limited. Specific examples of the substituents include hydroxy, the above-mentioned alkoxy, carboxy, the above-mentioned alkoxycarbonyl, the above-mentioned acyloxy, and amino which may be substituted by the above-mentioned alkyl, the above-mentioned arralkyl or the above-mentioned amino-protecting group. Preferred are hydroxy, alkoxy, carboxy, alkoxycarbonyl and acyloxy.

"Optionally substituted" of "optionally substituted cycloalkyl", "optionally substituted aryl" and "optionally substituted heterocyclic group having one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom" at R<sup>12</sup> means that they may have 1 to 3 substituent(s) on the ring wherein said substituents may be the same or different. The position of the substituent(s) is optional and is not particularly limited. Specific examples of the substituents include hydroxy, the above-mentioned lower alkoxy, mercapto, the above-mentioned lower alkylthio, carboxy, the above-mentioned lower alkyl, the above-mentioned lower alkyl, amino which may be substituted by the above-mentioned lower alkyl, the above-mentioned halogen atom, carbamoyl, cyano, the above-mentioned acyl, nitro, sulfamoyl, alkoxythiocarbonyl, thioalkanoyl, alkylsulfonyl such as methylsulfonyl and ethylsulfonyl, azomethine which may be substituted by the above-mentioned lower alkyl, the above-mentioned aryl or the above-mentioned aryl or the above-mentioned lower alkyl, aminooxy which may be substituted by the above-mentioned lower alkyl, the above-mentioned aryl or the above-mentioned aralkyl, and alkylsulfinyl such as methylsulfinyl. Preferred are hydroxy, lower alkyl, halogen atom, lower alkoxy, amino and carboxy.

"Optionally substituted" of "optionally substituted aralkyl" at R<sup>12</sup> means that it may have 1 to 3 substituent(s) on aryl wherein said substituents may be the same or different. The position of the substituent(s) is optional and is not particularly limited. Specific examples of the substituents include the above-mentioned lower alkyl, the above-mentioned lower alkyl, the above-mentioned lower alkyl or the above-mentioned acyl, the above-mentioned alkoxycarbonyl, the above-mentioned aryloxycarbonyl, the above-mentioned aryloxy, the above-mentioned alkylthio, the above-mentioned aryloxy, the above-mentioned aryloxy. It is above-mentioned aryloxy and the above-mentioned aryloxy. It is above-mentioned aryloxy. It is above-mentioned aryloxy and the above-mentioned aryloxy. It is above-mentioned aryloxy and halogen atom.

The compounds of the present invention which is shown by the formula (I) can be synthesized by, for example, the following method, to which the synthesis method of the compounds of the present invention is not limited.

wherein

R'

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which is more specifically protected  $R_a$ , protected alkoxy substituted by  $R_a$ , protected alkylthio substituted by  $R_a$ , protected alkylthio substituted by  $R_a$ , protected alkylamino substituted by  $R_a$ , protected and optionally substituted non-aromatic heterocyclic group containing nitrogen, or protected hydroxy, wherein when R is dimethylamino, N-methylpiperazinyl or N-methylpiperidyl, R means R itself, since R does not need to be protected, wherein  $R_a$  is amino, guanidino, amidino, carbamoyl, ureido, thioureido, hydrazino, hydrazinocarbonyl or imino, these groups being optionally substituted by a substituent selected from the group consisting of lower alkyl, halogenated lower alkyl, cycloalkyl, aralkyl, aryl and amino-pro-

is R protected by hydroxy-protecting group or amino-protecting group,

tecting group;

R<sup>14</sup> is carboxy-protecting group such as

is carboxy-protecting group such as methyl, ethyl, tert-butyl, allyl, phenyl, benzyl, trichloroethyl, p-nitrobenzyl, trimethylsilyl, tert-butyldimethylsilyl,

methoxymethyl and 2-trimethylsilylethyl;

W is halogen atom;

A' is A without one end methylene;

is hydrogen atom or substituent which activates X such as triphenylphos-

phonium, triphenylphosphonate and arylsulfonyl; and

A, X, M, m, R, R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup>, R<sup>4</sup>, R<sup>5</sup>, R<sup>6</sup> and R<sup>7</sup> are as defined above.

(Step 1)

Z

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The compound (VI) can be synthesized by reacting compound (II) and compound (III) in the presence of a combined condensing agent of triphenylphosphine, trimethylphosphine, triethylphosphine, triphenyl phosphite, trimethylphosphite, triethyl phosphite, and the like, and diisopropyl azodicarboxylate, diethyl azodicarboxylate, dicyclohexyl azodicarboxylate, and the like, in an organic solvent such as ether, tetrahydrofuran, dioxane, dichloromethane, chloroform, benzene, toluene and dimethylformamide, or a mixed solvent thereof, under ice-cooling to under heating.

This method is particularly preferable when X is oxygen atom or sulfur atom.

The compound (VI) can be also synthesized by the following method.

(Step 2)

The compound (VI) can be synthesized by reacting compound (IV) and compound (III) in the presence of a base such as sodium hydride, potassium hydride, lithium hydride, potassium carbonate, sodium carbonate, potassium tert-butoxide, lithium diisopropylamide, methyllithium, n-butyllithium, sec-butyllithium and tert-butyllithium, in an organic solvent such as dimethylformamide, methylene chloride, tetrahydrofuran, ether, benzene and toluene, or a mixed solvent thereof, at a temperature of from -78°C to under heating.

This method is particularly preferable when X is sulfur atom or oxygen atom.

When X is -SO- or -SO<sub>2</sub>-, the corresponding sulfide obtained in the above Step 1 or Step 2 is oxidized with an oxidizing agent such as hydrogen peroxide, peracetic acid, metaperiodate, metachloroperbenzoic acid, acyl nitrate and dinitrogen tetraoxide, to synthesize compound (VI).

The compound (VI) wherein X is particularly -NR8- or -CR9R10- can be also synthesized by the following method.

(Step 3)

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The compound (VI) can be synthesized by condensing compound (V) and compound (III) in the presence of a suitable base (e.g., lithium diisopropylamide, lithium bis(trimethylsilyl)amide, potassium bis(trimethylsilyl)amide, n-butyllithium, potassium tert-butoxide, sodium hydroxide, potassium hydroxide, sodium hydride and potassium hydride) as necessary, in water or an organic solvent such as methanol, ethanol, dimethylformamide, ether, dioxane, tetrahydrofuran, ethyl acetate, diisopropyl ether, dimethoxyethane, toluene, hexane and dimethyl sulfoxide, or a mixed solvent thereof, and subjecting the obtained compound to catalytic reduction using hydrogen gas in the presence of a metallic catalyst such as platinum black, platinum oxide, palladium black, palladium oxide, palladium hydroxide, palladium carbon and Raney nickel, or treating the compound with a reducing agent such as sodium borohydride, sodium cyanoborohydride, trimethylsilane, triethylsilane, alkali metal-ammonia, alkali metal-ethylamine, sodium amalgam and potassium amalgam.

The compound (I) can be synthesized by subjecting compound (VI) obtained in the above Step 1, 2 or 3 to the following Steps 4-6.

(Step 4)

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The compound (VII) can be synthesized by reacting compound (VI) in the presence of a hydroxide or carbonate of alkali metal such as sodium, potassium and lithium, or a base such as 1,5-diazabicyclo[4.3.0]non-5-ene and 1,8-diazabicyclo[5.4.0]undec-7-ene, or an acid such as hydrochloric acid, hydrobromic acid, sulfuric acid, hydrogen chloride, hydrogen bromide, hydrogen fluoride, acetic acid and trifluoroacetic acid, in water or an organic solvent such as methanol, ethanol, dichloromethane, chloroform, tetrahydrofuran, toluene and xylene or a mixed solvent thereof, under ice-cooling to under heating, or by subjecting compound (VI) to catalytic reduction using hydrogen gas in an organic solvent such as methanol, ethanol, dimethylformamide, ether, dioxane, tetrahydrofuran and acetic acid or a mixed solvent thereof, in the presence of a metallic catalyst such as platinum black, platinum oxide, palladium black, palladium

oxide, palladium carbon and Raney nickel, or by reacting compound (VI) in the presenc of quaternary ammonium fluoride such as tetraethylammonium fluoride and tetra-n-butylammonium fluoride, in an organic solvent such as tetrahydrofuran, dimethylformamide and dimethyl sulfoxide or a mixed solvent thereof, under ice-cooling to under heating.

#### 5 (Step 5)

The compound (I') can be synthesized by reacting compound (VII) and compound (VIII) using a condensing agent such as 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide hydrochloride (WSC • HCI), dicyclohexylcarbodiimide (DCC), diphenylphosphoryl azide (DPPA) and carbonyldiimidazole (CDI), in the presence of an activating agent such as 1-hydroxybenzotriazole (HOBT), hydroxysuccinimide (HOSu) and N-hydroxy-5-norbornene-2,3-dicarboximide (HONB) as necessary, in an organic solvent such as dimethylformamide, dichloromethane, chloroform, acetonitrile, tetrahydrofuran, dimethyl sulfoxide, carbon tetrachloride and toluene or a mixed solvent thereof, under ice-cooling to under heating. When compound (VIII) is, for example, hydrochloride, this reaction can be carried out in the presence of a base such as triethylamine, N-methylmorpholine and 4-dimethylaminopyridine. When R<sup>7</sup> is a group having hydroxy, such as -CONHOH and -CH<sub>2</sub>OH, compound (VIII) wherein said hydroxy is protected in advance is used.

(Step 6)

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This step aims at eliminating the hydroxy-protecting group or amino-protecting group at R', and can be carried out according to a suitable known method. For example, when the amino-protecting group at R' is Boc (tert-butoxycarbonyl group), compound (I') is reacted in the presence of an acid such as hydrochloric acid, hydrobromic acid, trifluoroacetic acid, p-toluenesulfonic acid, methanesulfonic acid, hydrogen chloride-dioxane, hydrogen chloride-ether and hydrogen chloride-ethyl acetate, in water or an organic solvent such as dioxane, ether, dichloromethane, tetrahydrofuran, methanol, ethanol, chloroform, benzene, toluene and ethyl acetate or a mixed solvent thereof or without solvent, to give compound (I). When the amino protecting group is, for example, benzyloxycarbonyl group, compound (I) can be synthesized by catalytic hydrogenation using hydrogen gas in water or an organic solvent such as methanol, ethanol, dimethylformamide, ether, dioxane, tetrahydrofuran and acetic acid or a mixed solvent thereof, in the presence of a metallic catalyst such as palladium carbon, platinum oxide and Raney nickel. When R' is hydroxy protected by hydroxy-protecting group, compound (I) can be synthesized by a conventional method such as catalytic hydrogenation. When R' is protected at hydroxy, the hydroxy-protecting group is eliminated by a conventional method such as catalytic hydrogenation, and thereafter or simultaneously therewith, the above Step is carried out.

The compound (I) wherein R<sup>7</sup> is carboxyl group can be synthesized by, for example, subjecting compound (I') wherein R<sup>7</sup> is tert-butoxycarbonyl group or benzyloxycarbonyl group to the above-mentioned reaction.

55 wherein

W<sup>1</sup> is -COW<sup>3</sup>, -SO<sub>2</sub>W<sup>3</sup> or -O-COW<sup>3</sup> wherein W<sup>3</sup> is hydroxy or halogen atom; W<sup>2</sup> is hydroxy, mercapto or -NR<sup>8</sup>H wherein R<sup>8</sup> is as defined above; and A, X, M, R', R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup>, R<sup>4</sup> and R<sup>14</sup> are as defined above.

The compound (VI) wherein X is -COO-, -CONR<sup>8</sup>-, -SO<sub>2</sub>NR<sup>8</sup>-, -COS-, -OOC-NR<sup>8</sup>- or -O-CO-O- can be also synthesized by the following method.

(Step 7)

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The compound (VI) can be synthesized by reacting compound (IX) and compound (X) using a condensing agent such as WSC • HCI, DCC, DPPA and CDI, in the presence of an activating agent such as HOBT, HOSu and HONB as necessary, in an organic solvent such as dimethylformamide, dichloromethane, chloroform, acetonitrile, tetrahydro-turan, dimethyl sulfoxide, carbon tetrachloride and toluene or a mixed solvent thereof, under ice-cooling to under heating (this reaction can be carried out in the presence of a base such as triethylamine, N-methylmorpholine, pyridine, 4-dimethylaminopyridine and N-methylpiperidine), or in the presence of a hydroxide or carbonate of alkali metal such as sodium, potassium and lithium, or a base such as triethylamine, pyridine, N-methylmorpholine, N-methylpiperidine and 4-dimethylaminopyridine, in water or an organic solvent such as dimethylformamide, dichloromethane, chloroform, tetrahydrofuran, dimethyl sulfoxide, benzene and toluene or a mixed solvent thereof, under ice-cooling to under heating.

The compound (VI) wherein X is -OOC-, -NR<sup>8</sup>CO-, -NR<sup>8</sup>SO<sub>2</sub>- or -NR<sup>8</sup>-COO- can be also synthesized by the following method.

20 (Step 8)

The compound (VI) can be synthesized using compound (XI) and compound (XII) according to the method of the above-mentioned Step 7.

When X is a divalent aromatic heterocyclic group having one or more hetero atoms selected from nitrogen atom, sulfur atom and oxygen atom, such as divalent oxadiazole ring, compound (VI) can be also synthesized by the following method.

(Step 10) 
$$R'-A$$
  $N$   $R^1$   $R^2$   $R^3$   $R^4$   $R^4$   $(VI')$ 

wherein A, M, R', R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup>, R<sup>4</sup> and R<sup>14</sup> are as defined above.

50 (Step 9)

The compound (XIV) can be synthesized by reacting compound (XIII) and compound (XIV) using a condensing agent such as WSC • HCI, DCC, DPPA and CDI, in the presence of an activating agent such as HOBT, HOSu and HONB as necessary, in an organic solvent such as dimethylformamide, dichloromethane, chloroform, acetonitrile, tetrahydrofuran, dimethyl sulfoxide, carbon tetrachloride and toluene or a mixed solvent thereof, under ice-cooling to under heating. This reaction can be carried out in the presence of a base such as trimethylamine, N-methylmorpholine, pyridine, 4-dimethylaminopyridine and N-methylpiperidin.

(Step 10)

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The compound (VI') can be synthesized by heating compound (XV) in an organic solvent such as toluene, dioxane, tetrahydrofuran, benzene and xylene, or a mixed solvent thereof.

The compound (I) can be synthesized by treating compound (VI) and compound (VI) obtained in the above Steps 7, 8 and 10 by the same method as in the above Steps 4-6.

When at least one of R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup> and R<sup>4</sup> of compound (I) is a halogen atom, compound (I) can be also synthesized by the following method.

wherein

R1', R2', R3' and R4'

are the same or different and each is hydrogen atom, hydroxy, alkoxy, mercapto, alkylthio, nitro, cyano, carboxy, alkoxycarbonyl, aryloxycarbonyl, acyl, alkyl which may be substituted by a substituent selected from hydroxy, lower alkoxy and halogen atom, amino which may be substituted by a substituent selected from alkyl, aryl, aralkyl and amino-protecting group, or -O-CO-R<sup>11</sup>

wherein R11 is as defined above,

provided that at least one of them is hydrogen atom; and

R<sup>7</sup> are as defined above.

A, X, M, m, R',  $R^1$ ,  $R^2$ ,  $R^3$ ,  $R^4$ ,  $R^5$ ,  $R^6$  and  $R^7$ 

(Step 11)

The compound (I') can be synthesized by reacting compound (I'') in the presence of a halogenating agent such as tert-butyl hypochlorite, tert-butyl hypobromite, tert-butyl hypoiodite, sulfuryl chloride, sulfuryl bromide, thionyl chloride, thionyl bromide, fluorine, chlorine, bromine, iodine, hydrogen fluoride, silver difluoride and xenon difluoride, in an organic solvent such as dichloromethane, chloroform, acetonitrile, toluene, benzene, ether, tetrahydrofuran, dioxane, methanol, ethanol, carbon tetrachloride and ethyl acetate, or a mixed solvent thereof, or without solvent, under ice-cooling to under heating. When the protective group is removed by this step, a re-protection is applied. In the case of Boc, for example, the compound is protected with di-tert-butyl dicarbonate and the like in the presence of a suitable base such as triethylamine and pyridine.

The compound (I) can be synthesized by treating the obtained compound (I') by the same method as in the above Step 6.

The above Step 11 may be carried out after synthesizing compound (VI) corresponding to compound (I"). The subsequent same treatment as in the above Steps 4-6 gives compound (I).

The compound (I) wherein at least one of  $R^1$ ,  $R^2$ ,  $R^3$  and  $R^4$  is -O-CO- $R^{11}$  can be also synthesized by the following method.

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wherein

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R1", R2", R3" and R4"

are the same or different and each is hydrogen atom, hydroxy, halogen atom, alkoxy, mercapto, alkylthio, nitro, cyano, carboxy, alkoxycarbonyl, aryloxycarbonyl, acyl, alkyl which may be substituted by a substitutent selected from hydroxy, lower alkoxy and halogen atom, or amino which may be substituted by a substituent selected from alkyl, aryl, aralkyl and amino-protecting group, wherein at least one of them is hydroxy; and

A, X, M, m, W, R', R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup>, R<sup>4</sup>, R<sup>5</sup>, R<sup>6</sup>, R<sup>7</sup> and R<sup>11</sup> are as defined above.

(Step 12)

The compound (I') can be synthesized by reacting compound (I") with compound (XVI) in an organic solvent such as dichloromethane, chloroform, ether, tetrahydrofuran, dioxane, benzene, toluene, dimethylformamide, ethyl acetate and acetonitrile or a mixed solvent thereof, in the presence of a base such as pyridine, triethylamine, N-methylmorpholine, N-methylpiperidine and 4-dimethylaminopyridine.

The compound (I) can be synthesized by reacting the obtained compound (I') by the same method as in the above Step 6.

The compound of the formula (I) of the present invention can be also synthesized by the following synthetic method.

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$$Z-X$$
  $M$   $COOH$   $+$   $R^{7}$   $R^{6}$   $R^{1}$   $R^{2}$   $CH_{2}$   $M$   $CON$   $R^{7}$   $R^{5}$   $R^{3}$   $R^{4}$   $R^{5}$   $R^{5}$   $R^{5}$   $R^{1}$   $R^{2}$   $R^{4}$   $R^{5}$   $R^{5}$   $R^{5}$   $R^{5}$   $R^{5}$   $R^{5}$   $R^{5}$   $R^{7}$   $R^{1}$   $R^{2}$   $R^{5}$   $R^{5}$   $R^{5}$   $R^{7}$   $R^{1}$   $R^{2}$   $R^{5}$   $R^{5}$   $R^{5}$   $R^{1}$   $R^{2}$   $R^{5}$   $R^{5}$   $R^{5}$   $R^{5}$   $R^{7}$   $R^{1}$   $R^{2}$   $R^{5}$   $R^{5}$   $R^{5}$   $R^{7}$   $R^{1}$   $R^{2}$   $R^{5}$   $R^{5}$   $R^{5}$   $R^{5}$   $R^{7}$   $R^{1}$   $R^{2}$   $R^{5}$   $R$ 

wherein A, A', X, M, m, W, Z, R',  $R^1$ ,  $R^2$ ,  $R^3$ ,  $R^4$ ,  $R^5$ ,  $R^6$  and  $R^7$  are as defined above.

(Step 13)

The compound (XVII) can be synthesized by subjecting compound (III') and compound (VIII) to the same reaction as in the above Step 5.

(Step 14)

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The compound (I') can be synthesized by subjecting compound (II) and compound (XVII) to the same reaction as in the above Step 1.

The compound (I') can be also synthesized by the following method.

(Step 15)

The compound (I') can be synthesized by subjecting compound (IV) and compound (XVII) to the same reaction as in the above Step 2.

The compound (i') wherein X is -NR8- or -CR9R10- can be also synthesized by the following method.

(Step 16)

The compound (I') can be synthesized by subjecting compound (V) and compound (XVII) to the same reaction as in the above Step 3.

The compound (I) can be synthesized by subjecting compound (I') obtained in the above Steps 14-16 to the same reaction as in the above Step 6.

The compound (I') wherein X is -COO-, -CONR<sup>8</sup>-, SO<sub>2</sub>NR<sup>8</sup>-, -COS-, -OOC-NR<sup>8</sup>- or -O-CO-O- can be also synthesized by the following method.

wherein A, X, M, m, W1, W2, R1, R1, R2, R3, R4, R5, R6 and R7 are as defined above.

(Step 17)

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The compound (XVIII) can be synthesized by subjecting compound (X') and compound (VIII) to the same reaction as in the above Step 5.

(Step 18)

The compound (I') can be synthesized by subjecting compound (IX) and compound (XVIII) to the same reaction as in the above Step 7.

The compound (I') wherein X is -OOC-, -NR8CO-, -NR8CO-, -NR8CO- or -NR8-COO- can be also synthesized by the follow-

ing method.

wherein A, X, M, m, W<sup>1</sup>, W<sup>2</sup>, R', R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup>, R<sup>4</sup>, R<sup>5</sup>, R<sup>6</sup> and R<sup>7</sup> are as defined above.

(Step 19)

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The compound (XIX) can be synthesized by subjecting compound (XII') and compound (VIII) to the same reaction as in the above Step 5.

(Step 20)

The compound (I') can be synthesized by subjecting compound (XI) and compound (XIX) to the same reaction as in the above Step 8.

The compound (I) can be synthesized by subjecting compound (I') obtained in the above Step 18 and Step 20 to the same reaction as in the above Step 6.

When X is -CR9R10-, -CO-, -C=C- or -CS-, the following step can be used for synthesis.

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wherein A, A', M, X, m, W<sup>1</sup>, W<sup>2</sup>, R', R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup>, R<sup>4</sup>, R<sup>5</sup>, R<sup>6</sup>, R<sup>7</sup> and R<sup>14</sup> are as defined above.

## 30 (Step 21)

The compound (XXI) can be synthesized by reacting the corresponding Grignard reagent (IV') obtained from compound (IV) by a conventional method, with compound (XX) in an organic solvent such as ether, tetrahydrofuran and dioxane or a mixed solvent thereof, at a temperature of from -78°C to under heating.

## (Step 22)

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The compound (VI") can be synthesized by reacting compound (XXI) in the presence of an oxidizing agent such as chromic anhydride, pyridinium chlorochromate, manganese dioxide, sodium hypochlorite and ruthenium tetraoxide, in an organic solvent such as ether, tetrahydrofuran and dioxane or a mixed solvent thereof, under ice-cooling to under heating.

The compound (VI") wherein X is -CS- can be synthesized by reacting compound (VI") obtained by the above method, in the presence of hydrogen sulfide, phosphorus pentasulfide, 2,4-bis(4-methoxyphenyl)-1,3-dithia-2,4-diphosphetane-2,4-disulfide (Lawsson's reagent) and the like, in an organic solvent such as benzene, toluene, methanol and ethanol or a mixed solvent thereof, under ice-cooling to under heating.

# (Step 23)

The compound (VI") can be synthesized by reacting compound (XXI) in the presence of a reducing agent such as triethylsilane, lithium alminium hydride-alminium chloride, sodium borohydride-trifluoroborane, sodium cyanoborohydride-methyl iodide and triphenylphosphonium, in an organic solvent such as ether, tetrahydrofuran and dioxane, or a mixed solvent thereof, at a temperature of from -78°C to under heating.

## (Step 24)

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The compound (VI<sup>\*\*\*</sup>) can be synthesized by reacting compound (XXI) in the presence of sulfuric acid, phosphoric acid, potassium hydrogensulfate, oxalic acid, p-toluenesulfonic acid, boron trifluoride-etherate, thionyl chloride-pyridine, phosphorus oxychloride-pyridine, methanesulfonyl chloride-pyridine, p-toluenesulfonyl chloride-pyridine, and the like, in

an organic solvent such as ether, tetrahydrofuran and dioxane, or a mixed solvent thereof, under ice-cooling to under heating.

The compound (I) can be synthesized by treating compound (VI"), (VI"') or (VI"'') obtained in the above Steps 22-24 by the same method as in the above Steps 4-6.

The compound of the formula (I) can be converted to a pharmaceutically acceptable acid addition salt by a conventional method by treating same with an inorganic acid (e.g., hydrochloric acid, sulfuric acid, phosphoric acid, hydrobromic acid and nitric acid) or organic acid (e.g., oxalic acid, maleic acid, humaric acid, malic acid, tartaric acid, succinic acid, citric acid, acetic acid, lactic acid, methanesulfonic acid, p-toluenesulfonic acid, benzoic acid, valeric acid, malonic acid, nicotinic acid and propionic acid).

The compound thus obtained can be separated and purified by a known method for separation and purification, such as concentration, concentration under reduced pressure, solvent extraction, precipitation, recrystallization and chromatography.

The compound of the present invention includes one or more stereoisomers due to an asymmetric carbon, and such isomers and mixtures thereof are also encompassed in the present invention. In addition, hydrates and solvates with pharmaceutically acceptable organic solvents, as well as prodrugs of the compound of the present invention are also encompassed in the present invention.

The compound of the present invention shows superior effects of suppressing production of inflammatory cytokines in mammals such as human, rabbit, dog and cat, and is useful for the prophylaxis and treatment of noninfectious or infectious diseases accompanied by neutrophile infiltration, which are represented by rheumatic diseases (e.g., rheumatoid arthritis); arthritis due to gout; systemic lupus erythematosus; dermatopathy (e.g., psoriasis, pustulosis and atopic dermatitis); respiratory diseases (e.g., bronchial asthma, bronchitis, ARDS and diffused interstitial pneumonia); inflammatory enteric diseases (e.g., ulcerative colitis and Crohn's disease); acute or chronic hepatitis inclusive of fulminant hepatitis; acute or chronic glomerulonephritis; nephropyelitis; uveitis caused by Behcet disease and vogt-Koyanagi Harada disease; Mediterranean fever (polyserositis); ischemic diseases (e.g., myocardial infarction); and systemic circulatory failure and multi-organ failure caused by sepsis.

The suppressive effect of the compound of the present invention on inflammatory cytokines such as IL-6 and GM-CSF has been also acknowledged.

When the compound of the formula (I) of the present invention or a pharmaceutically acceptable salt thereof is used as a pharmaceutical preparation comprising same as an active ingredient, it is generally admixed with a pharmaceutically acceptable carrier, excipient, diluent, extender, disintegrator, stabilizer, preservative, buffer, emulsifying agent, aromatic, coloring, sweetener, thickener, flavor, solubilizer and other additives such as water, vegetable oil, alcohols (e.g., ethanol and benzyl alcohol), polyethylene glycol, glycerol triacetate, gelatin, lactose and carbohydrate (e.g., starch), magnesium stearate, talc, lanolin, white petrolatum known *per se* to give a pharmaceutical composition in the form of tablet, pill, powder, granule, suppository, injection, eye drop, liquid, capsule, troche, aerosol, elixir, suspension, emulsion, syrup and the like, which is administered orally or parenterally.

While the dose varies depending on the kind and severity of diseases, compound to be administered, administration route, age, sex, body weight etc. of the patients, and so on, when it is orally administered to an adult patient, for example, the daily dose is generally 0.01 - 1,000 mg, preferably about 0.1 - 100 mg, and when it is intravenously administered to an adult patient, for example, the daily dose is generally 0.01 - 1,000 mg, preferably about 0.05 - 50 mg, which is administered in one or several doses.

The present invention is described in more detail by illustrative Preparative Examples and Examples, to which the present invention is not limited.

Hereunder follow Preparative Examples of the intermediate compounds shown in Table 1.

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Table 1

5	Preparative Example		Preparative Example	
10	1	Me OH HO — COOH C1	4	H <sub>2</sub> N
15	2	H <sub>2</sub> N OH COOMe	. 5	H <sub>2</sub> N
20	3	Boc - N COOH	6	H <sub>2</sub> N
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Preparative Example	
4	Ph H₂N CONH-Ph • HC1
5	Ph H <sub>2</sub> N CONHO Ph • HC1
6	H <sub>2</sub> N N Me • HCl
7	H <sub>2</sub> N

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## Preparative Example 1

## 5-Chloro-2,4-dihydroxy-3-methylbenzoic acid

To a solution of 2,4-dihydroxy-3-methylbenzoic acid methyl ester (9.9 g) in ethyl acetate (100 ml) was added tert-butyl hypochlorite (12.3 ml) under ice-cooling. After stirring for 2 hours, hexane (200 ml) was added, and the mixture was cooled with ice to allow precipitation of crystals. The crystals were collected by filtration, and dissolved in a mixed solvent of methanol (20 ml) and tetrahydrofuran (THF, 20 ml). A 1M lithium hydroxide solution (40 ml) was added to the solution, and the mixture was refluxed under heating for 18 hours. The reaction mixture was concentrated, and a 10% aqueous citric acid solution was added to the residue, which was followed by extraction with ethyl acetate. The organic layer was washed with water and saturated brine, and dried over anhydrous sodium sulfate. The organic layer was concentrated under reduced pressure to give the title compound (4.14 g, yield 37%).

Preparative Example 2

## Methyl 2-hydroxybenzoate-4-carboxamide oxime

To a solution of 2-hydroxy-4-cyanobenzoic acid methyl ester (2.00 g) in methanol (30 ml) were added water (6 ml), hydroxylamine hydrochloride (1.57 g) and sodium hydrogencarbonate (1.9 g), and the mixture was stirred with heating at 70°C for 3 hours. The reaction mixture was concentrated, diluted with a 10% aqueous citric acid solution, and filtered. The filtrate was concentrated under reduced pressure, and the residue was purified by silica gel column chromatography (developing solvent: hexane/ethyl acetate=3/2 v/v) to give the title compound (823 mg, yield 35%).

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#### Preparative Example 3

1-tert-Butoxycarbonyl-4-ethylisonipecotic acid

## (1) 1-tert-Butoxycarbonyl-4-ethylisonipecotic acid ethyl ester

To a solution of 1-tert-butoxycarbonylisonipecotic acid ethyl ester (576 mg) in THF (15 ml) was added a solution of lithium diisopropylamide (290 mg) in THF (10 ml) in a stream of argon gas at -78°C, and the reaction mixture was stirred at the same temperature for 1 hour. Ethyl iodide (0.36 ml) was added to the above solution at -78°C, and the mixture was stirred for 18 hours. Water was added to the reaction mixture and the mixture was extracted with ethyl acetate. The organic layer was washed successively with 1N hydrochloric acid, water and saturated brine, and dried over anhydrous magnesium sulfate. The organic layer was concentrated under reduced pressure to give the title compound (585 mg, yield 92%).

## 15 (2) 1-tert-Butoxycarbonyl-4-ethylisonipecotic acid

To a solution of 1-tert-butoxycarbonyl-4-ethylisonipecotic acid ethyl ester (570 mg) in ethanol (10 ml) was added a 1M lithium hydroxide solution (8 ml), and the mixture was refluxed under heating for 20 hours. Then, the reaction mixture was concentrated, and water was added to the residue. The aqueous layer was washed with ether, acidified with 1N hydrochloric acid, and extracted with ether. The organic layer was dried over anhydrous magnesium sulfate, and concentrated under reduced pressure to give the title compound (233 mg, yield 45%).

## Preparative Example 4

#### 25 L-Phenylalanylaminobenzene hydrochloride

To a solution of N-tert-butoxycarbonyl-L-phenylalanine hydrochloride (2.65 g) and aniline (1.02 g) in dimethylformamide (DMF, 50 ml) were added 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide hydrochloride (WSC • HCl) and hydroxybenzotriazole (HOBT, 1.5 g) at room temperature, and the mixture was stirred for 6 hours. Water was added to the reaction mixture and the mixture was extracted with ethyl acetate. The organic layer was washed successively with a 10% aqueous citric acid solution, water, a saturated aqueous sodium hydrogencarbonate solution, water and saturated brine, and dried over anhydrous magnesium sulfate. The solvent was distilled away under reduced pressure to give N-tert-butoxycarbonyl-L-phenylalanylaminobenzene. To a solution of the obtained compound in dichloromethane (20 ml) was added trifluoroacetic acid (10 ml) at room temperature, and the mixture was stirred for 1 hour. Toluene (10 ml) was added to the reaction mixture, and the mixture was concentrated under reduced pressure. A 1M hydrogen chlorideether solution (10 ml) was added to the residue, and crystallization gave the title compound (1.45 g, yield 52%).

## Preparative Example 5

# 40 L-Phenylalanyl-O-benzylhydroxyamide hydrochloride

The title compound (2.48 g, yield 92%) was obtained in the same manner as in Preparative Example 4 above, using N-tert-butoxycarbonyl-L-phenylalanine (2.65 g) and O-benzylhydroxylamine hydrochloride (1.60 g).

## 45 Preparative Example 6

# 1-(3-Methyl-1,2,4-oxadiazol-5-yl)-2-phenylethylamine hydrochloride

To a solution of acetamide oxime [2.67 g, J. Saunders et al., J. Med. Chem., 33, 1128 (1990)] in THF (125 ml) was added 60% sodium hydride (1.44 g) in oil, and the mixture was refluxed under heating for 1 hour. Then, the reaction mixture was allowed to cool, and a solution of N-tert-butoxycarbonyl-L-phenylalanine methyl ester (8.38 g) in THF (40 ml) was added at room temperature. The mixture was refluxed under heating for 20 minutes. The mixture was allowed to cool, and water (10 ml) was added, which was followed by concentration under reduced pressure. A 10% aqueous citric acid solution was added to the residue, and the mixture was extracted with ethyl acetate. The organic layer was washed with saturated brine, dried over anhydrous sodium sulfate, and concentrated under reduced pressure. The residue was purified by silica gel column chromatography (developing solvent: hexane/ethyl acetate=3/1 v/v) to give 4.43 g of N-tert-butoxycarbonyl-1-(3-methyl-1,2,4-oxadiazol-5-yl)-2-phenylethylamine. This compound was added to a 4N hydrogen chloride-dioxane solution (50 ml), and the mixture was stirred at room temperature for 2 hours. Toluene was

added to the reaction mixture, and the mixture was concentrated under reduced pressur. Ether was added to the residue for crystallization to give the title compound (3.25 g, yield 47%).

#### Preparative Example 7

O-Benzyl-L-phenylalaninol

To a solution of L-phenylalaninol (11.78 g) in THF (200 ml) was gradually added 60% sodium hydride (3.43 g) in oil at room temperature. Twenty minutes later, the reaction mixture was refluxed under heating for 1 hour. Then, the mixture was allowed to cool, followed by gradual addition of benzyl bromide (9.27 ml) under ice-cooling, and stirred at room temperature for 16 hours. The reaction mixture was added to saturated brine, and extracted with ether. The organic layer was extracted with 10% hydrochloric acid. The aqueous layer was made alkaline with an aqueous sodium hydroxide solution, and extracted with ether. The organic layer was washed with saturated brine, dried over anhydrous magnesium sulfate, and concentrated under reduced pressure to give the title compound (14.5 g, yield 77%).

## Example 1

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N-[3,5-Dichloro-2-hydroxy-4-(4-methylaminobutoxy)benzoyl]-L-phenylalanine methyl ester hydrochloride

20 Step 1) 3,5-Dichloro-2-hydroxy-4-(4-tert-butoxycarbonylmethylaminobutoxy)benzoic acid methyl ester (VI)

To a solution of 4-tert-butoxycarbonylmethylamino-1-butanol (3 g) and known 3,5-dichloro-2,4-dihydroxybenzoic acid methyl ester (3.85 g) in THF (80 ml) were added triphenylphosphine (4.26 g) and diisopropyl azodicarboxylate (3.2 ml) under ice-cooling, and the mixture was stirred for 16 hours. The reaction mixture was concentrated under reduced pressure, and the residue was purified by silica gel column chromatography (developing solvent: hexane/ethyl acetate=4/1 v/v) to give the title compound (5.2 g, yield 83%).

Step 4) 3,5-Dichloro-2-hydroxy-4-(4-tert-butoxycarbonylmethylaminobutoxy)benzoic acid (VII)

The compound (3.46 g) obtained in the above Step 1) was dissolved in a mixed solvent of methanol (12 ml)-THF (12 ml), and a 1M lithium hydroxide solution (24 ml) was added to the mixture, which was followed by stirring with heating at 60°C for 2 hours. After cooling with ice, the mixture was concentrated under reduced pressure. A 10% aqueous citric acid solution (50 ml) was added to the residue to acidify same, and the mixture was extracted with ether (50 ml). The organic layer was dried over anhydrous magnesium sulfate, and the solvent was distilled away under reduced pressure to give the title compound (3.22 g, yield 96%).

Step 5) N-[3,5-Dichloro-2-hydroxy-4-(4-tert-butoxycarbonylmethylaminobutoxy)benzoyl]-L-phenylalanine methyl ester (I')

To a solution of the compound (3 g) obtained in the above Step 4), L-phenylalanine methyl ester hydrochloride (1.59 g), WSC • HCl (1.41 g) and HOBT (1 g) in DMF (10 ml) was added dropwise triethylamine (1 ml) at room temperature, and the mixture was stirred for 14 hours. Water (60 ml) was added to the reaction mixture, and the mixture was extracted with ethyl acetate. The organic layer was washed successively with a 10% aqueous citric acid solution, water, a saturated aqueous sodium hydrogencarbonate solution, water and saturated brine, and dried over anhydrous sodium sulfate. Then, the solvent was distilled away under reduced pressure. The residue was purified by silica gel column chromatography (developing solvent: hexane/ethyl acetate=3/1 v/v) to give the title compound (2.72 g, yield 65%).

Step 6) N-[3,5-Dichloro-2-hydroxy-4-(4-methylaminobutoxy)benzoyl]-L-phenylalanine methyl ester hydrochloride (I)

To a solution of the compound (5 g) obtained in the above Step 5) in dioxane (10 ml) was added a 4N hydrogen chloride-dioxane solution (40 ml), and the mixture was stirred at room temperature for 1.5 hours. The reaction mixture was diluted with toluene, and concentrated under reduced pressure. Ether (50 ml) was added to the residue for crystallization to give the title compound (4.2 g, yield 95%, see Table 2).

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## Example 1'

N-[3,5-Dichloro-2-hydroxy-4-(4-methylaminobutoxy)benzoyl]-L-phenylalanin methyl ester hydrochloride

5 Step 13) N-(3,5-Dichloro-2,4-dihydroxybenzoyl)-L-phenylalanine methyl ester (XVII)

To a solution of 3,5-dichloro-2,4-dihydroxybenzoic acid (17 g), L-phenylalanine methyl ester hydrochloride (19.8 g), WSC • HCl (17.6 g) and HOBT (12.4 g) in DMF (70 ml) was added dropwise triethylamine (12.8 ml) at room temperature, and the mixture was stirred for 16 hours. Then, the mixture was post-treated in the same manner as in the above Example 1, Step 5) to give the title compound (18.32 g, yield 57%).

Step 14) N-[3,5-Dichloro-2-hydroxy-4-(4-tert-butoxycarbonylmethylaminobutoxy)benzoyl]-L-phenylalanine methyl ester (I')

To a solution of the compound (11.0 g) obtained in the above Step 13) and 4-tert-butoxycarbonylmethylamino-1-butanol (5.29 g) in THF (100 ml) were added triphenylphosphine (7.51 g) and diisopropyl azodicarboxylate (5.6 ml) under ice-cooling, and the mixture was stirred for 16 hours. The reaction mixture was concentrated under reduced pressure, and the residue was purified by silica gel column chromatography (developing solvent: hexane/ethyl acetate=3/1 v/v) to give the title compound (3.10 g, yield 21%).

Step 6) N-[3,5-Dichloro-2-hydroxy-4-(4-methylaminobutoxy)benzoyl]-L-phenylalanine methyl ester hydrochloride (I)

To a solution of the compound (10 g) obtained in the above Step 14) in dioxane (25 ml) was added dropwise a 4N hydrogen chloride-dioxane solution (88 ml) at room temperature. After 1.5 hours, toluene was added. The solvent was distilled away under reduced pressure, and ether (120 ml) was added to the residue for crystallization to give the title compound (8.4 g, yield 95%).

#### Example 2

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N-{3,5-Dichloro-2-hydroxy-4-{2-(4-methylpiperazin-1-yl)ethoxy|benzoyl}-L-phenylalanine ethyl ester dihydrochloride

Step 1) 3,5-Dichloro-2-hydroxy-4-[2-(4-methylpiperazin-1-yl)ethoxy]benzoic acid methyl ester (VI)

To a solution of 2-(4-methylpiperazin-1-yl)ethanol (14.42 g) and 3,5-dichloro-2,4-dihydroxybenzoic acid methyl ester (52.15 g) in chloroform (400 ml) were added triphenylphosphine (28.85 g) and azodicarboxylic acid diisopropyl (21.7 ml) at room temperature, and the mixture was stirred for 16 hours. 1N Hydrochloric acid (300 ml) was added to the reaction mixture for extraction to give a crude product of the title compound.

Step 4) 3,5-Dichloro-2-hydroxy-4-[2-(4-methylpiperazin-1-yl)ethoxy]benzoic acid (VII)

To the extract of the crude product obtained in the above Step 1) was added a 4M aqueous sodium hydroxide solution (125 ml), and the mixture was stirred under heating at 80°C for 2 hours. Acetic acid (12.3 g) was further added to the mixture. The mixture was stirred under ice-cooling, and applied to crystallization to give the title compound (27.880 g, yield 79%).

Step 5) N-{3,5-Dichloro-2-hydroxy-4-[2-(4-methylpiperazin-1-yl)ethoxy]benzoyl}-L-phenylalanine ethyl ester dihydrochloride (l'=l)

To a solution of the compound (958 mg) obtained in the above Step 4), L-phenylalanine ethyl ester hydrochloride (923 mg) and HOBT (445 mg) in acetonitrile (15 ml) was added WSC • HCI (632 mg) at room temperature, and the mixture was stirred for 25 hours. The reaction mixture was concentrated under reduced pressure, and chloroform was added to the residue. The mixture was washed successively with a saturated aqueous sodium hydrogencarbonate solution, water and saturated brine, dried over anhydrous sodium sulfate, and concentrated under reduced pressure. The residue was purified by silica gel column chromatography (developing solvent: chloroform/ methanol=10/1 v/v) to give a compound (1.386 g). Then, a 4N hydrogen chloride-ethyl acetate solution was added to a solution of the compound (1.003 g) in acetone (10 ml) for crystallization to give the title compound (1.073 g, yield 93%, see Table 2).

## Examples 3-87

The compounds of Example 3-87 were prepared in the same manner as in Example 1, Example 1' and Example 2 from the corresponding compounds (see Tables 3-45).

Example 88

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N-[2-Hydroxy-4-(4-methylaminobutyl)benzoyl]-L-phenylalanine methyl ester hydrochloride

Step 3) 4-[4-(tert-Butoxycarbonylmethylamino)butyl]-2-hydroxybenzoic acid methyl ester (VI)

(1) 4-[4-(tert-Butoxycarbonylmethylamino)-1-butenyl]-2-hydroxybenzoic acid methyl ester

To a solution of [(3-hydroxy-4-methoxycarbonyl)benzyl]triphenylphosphonium bromide (2.537 g), obtained by a known method, in THF (25 ml) was added dropwise a 2M lithium diisopropylamide-THF solution (5.5 ml) in a stream of argon at 0°C, and the mixture was stirred for 30 minutes. Then, a solution of 4-(tert-butoxycarbonylmethylamino) butylaldehyde (1.123 g), prepared by a known method, in THF (10 ml) was gradually added dropwise at 0°C, and the mixture was stirred at room temperature for 4 hours. A saturated aqueous ammonium chloride solution (1 ml) was gradually added, and the mixture was concentrated under reduced pressure, which was followed by extraction with toluene. The extract was washed with a 10% aqueous citric acid solution and saturated brine, dried over anhydrous sodium sulfate. and concentrated under reduced pressure. The residue was purified by silica gel column chromatography (developing solvent; hexane/ethyl acetate=4/1 v/v) to give the title compound (0.850 g, yield 51%).

(2) 4-[4-(tert-Butoxycarborrylmethylamino)butyl]-2-hydroxybenzoic acid methyl ester

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A solution of the compound (0.845 g) obtained in (1) above in methanol (20 ml) was vigorously stirred in the presence of 10% palladium-carbon (0.106 g) in a stream of hydrogen. After filtering through Celite, the mixture was concentrated under reduced pressure, and the residue was purified by silica gel column chromatography (developing solvent: hexane/ethyl acetate=4/1 v/v) to give the title compound (0.810 g, yield 95%).

Step 4) 4-[5-(tert-Butoxycarbonylmethylamino)butyl]-2-hydroxybenzoic acid (VII)

The compound (0.806 g) obtained in the above Step 3) was subjected to the same reaction as in the above Example 1, Step 4) to give the title compound (0.760 g, yield 98%).

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Step 5) N-[2-Hydroxy-4-(4-tert-butoxycarbonylmethylaminobutyl)benzoyl]-L-phenylalanine methyl ester (I')

The compound (0.753 g) obtained in the above Step 4) and L-phenylalanine methyl ester hydrochloride (0.552 g) was subjected to the same reaction as in the above Example 1, Step 5) to give the title compound (0.714 g, yield 63%).

Step 6) N-[2-Hydroxy-4-(4-methylaminobutyl)benzoyl)-L-phenylalanine methyl ester hydrochloride (I)

The compound (0.128 g) obtained in the above Step 5) was subjected to the same reaction as in the above Example 1, Step 6) to give the title compound (0.087 g, yield 78%, see Table 46).

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Examples 89, 90

The compounds of Examples 89 and 90 were prepared in the same manner as in Example 88 from the corresponding compounds (see Tables 46-47).

Example 91

N-[3,5-Dichloro-2-hydroxy-4-(5-methylaminopentyl)benzoyf]-L-phenylalanine methyl ester hydrochloride

Step 11) 4-[5-(tert-Butoxycarbonylmethylamino)pentyl]-3,5-dichloro-2-hydroxybenzoic acid methyl ester (VI)

To a solution of 4-[5-(tert-butoxycarbonylmethylamino)pentyl]-2-hydroxybenzoic acid methyl ester (3.95 g), obtained in the same manner as in the above Example 88, Step 3), in acetonitrile (35 ml) was added sulfuryl chloride

(9 ml) at room temperature, and the mixture was refluxed under h ating at 60°C for 1 hour. Toluene was added to the reaction mixture and the mixture was concentrated under reduced pressure. Dichloromethane (85 ml) was added to th residue. Then, triethylamine (7.85 ml) and di-tert-butyl dicarbonate (4.9 g) were added, and the mixture was stirred at room temperature for 1 hour. Water (50 ml) was added to the reaction mixture for washing, and the mixture was dried over anhydrous sodium sulfate, and concentrated under reduced pressure. The residue was purified by silica gel column chromatography (developing solvent:hexane/ethyl acetate=4/1 v/v) to give the titl compound (2.319 g, yield 50%).

Step 4) 4-[5-(tert-Butoxycarbonylmethylamino)pentyl]-3,5-dichloro-2-hydroxybenzoic acid (VII)

The compound (2.319 g) obtained in the above Step 11) was subjected to the same reaction as in the above Example 1, Step 4) to give the title compound (1.994 g, yield 89%).

Step 5) N-[4-(5-tert-Butoxycarbonylmethylaminopentyl)-3,5-dichloro-2-hydroxybenzoyl]-L-phenylalanine methyl ester (I')

The compound (2.874 g) obtained in the above Step 4) and L-phenylalanine methyl ester hydrochloride (1.522 g) was subjected to the same reaction as in the above Example 1, Step 5) to give the title compound (3.441 g, yield 86%).

Step 6) N-[3,5-Dichloro-2-hydroxy-4-(5-methylaminopentyl)benzoyl]-L-phenylalanine methyl ester hydrochloride (I)

The compound (3.426 g) obtained in the above Step 5) was subjected to the same reaction as in the above Example 1, Step 6) to give the title compound (2.525 g, yield 83%, see Table 48).

#### Examples 92-104

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The compounds of Examples 92-104 were prepared in the same manner as in Example 91 from the corresponding compounds (see Tables 48-54).

## Example 105

N-[2-Benzoyloxy-3,5-dichloro-4-(4-methylaminobutoxy)benzoyl]-L-phenylalanine methyl ester hydrochloride

Step 12) N-[2-Benzoyloxy-3,5-dichloro-4-(4-tert-butoxycarbonylmethylaminobutoxy)benzoyl]-L-phenylalanine methyl ester (I')

To a solution of the compound (212 mg), obtained in the above Example 1, Step 5), in dichloromethane (3 ml) were added pyridine (60 μl) and benzoyl chloride (80 μl) at room temperature, and the mixture was stirred for 30 minutes. Water (5 ml) was added to the reaction mixture and the mixture was extracted with ethyl acetate. The organic layer was washed successively with a 10% aqueous citric acid solution, a saturated aqueous sodium hydrogencarbonate solution, water and saturated brine, and dried over anhydrous magnesium sulfate. Then, the solvent was distilled away under reduced pressure. The residue was purified by silica gel column chromatography (developing solvent: hexane/ethyl acetate=3/1 v/v) to give the title compound (224 mg, yield 95%).

Step 6) N-[2-Benzoyloxy-3,5-dichloro-4-(4-methylaminobutoxy)benzoyl]-L-phenylalanine methyl ester hydrochloride (I)

The compound (224 mg) obtained in the above Step 12) was subjected to the same reaction as in the above Example 1, Step 6) to give the title compound (159 mg, yield 83%, see Table 55).

## Examples 106-125

The compounds of Examples 106-125 were prepared in the same manner as in Example 105 from the corresponding compounds (see Tables 55-65).

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## Example 126

N-[3,5-Dichloro-2-hydroxy-4-(4-methylaminobutoxy)benzoyf]-L-phenylalanine hydrochloride

5 Step 6) N-[3,5-Dichloro-2-hydroxy-4-(4-methylaminobutoxy)benzoyf]-L-phenylalanine hydrochloride (I)

To a solution of N-[3,5-dichloro-2-hydroxy-4-(4-tert-butoxycarbonylmethylaminobutoxy)benzoyl]-L-phenylalanine tert-butyl ester (490 mg), obtained in the same manner as in the above Example 1, Step 5), in dichloromethane (8 ml) was added trifluoroacetic acid (4 ml) at room temperature, and the mixture was stirred for 14 hours. Toluene was added to the reaction mixture and the mixture was concentrated under reduced pressure. A 1M hydrogen chloride-ether solution (5 ml) was added to the residue for crystallization to give the title compound (250 mg, yield 67%, see Table 66).

#### **Examples 127-135**

The compounds of Examples 127-135 were prepared in the same manner as in Example 126 from the corresponding compounds (see Tables 66-70).

# Example 136

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N-[3,5-Dichloro-2-hydroxy-4-(4-methylaminobutylamino)benzoyl]-L-phenylalanine methyl ester dihydrochloride

Step 16) N-[2-Hydroxy-4-(4-tert-butoxycarbonylmethylaminobutylamino)benzoyl]-L-phenylalanine methyl ester (I')

A solution of N-[(4-amino-2-hydroxy)benzoyl]-L-phenylalanine methyl ester (1.11 g) obtained in the same manner as in the above Example 1', Step 13) and 4-(tert-butoxycarbonylmethylamino)-1-butylaldehyde (711 mg) in methanol (20 ml) was stirred at room temperature in a stream of argon for 4 hours. 10% Palladium-carbon (200 mg) was added to the reaction mixture, and the mixture was subjected to catalytic hydrogenation using hydrogen gas under atmospheric pressure. Four hours later, the reaction mixture was filtered, and the filtrate was concentrated under reduced pressure. The residue was purified by silica gel column chromatography (developing solvent: hexane/ethyl acetate=3/2 v/v) to give the title compound (900 mg, yield 51%).

Step 11) N-[3,5-Dichloro-2-hydroxy-4-(4-tert-butoxycarbonylmethylaminobutylamino)benzoyl]-L-phenylalanine methyl ester (I')

To a solution of the compound (900 mg) obtained in the above Step 16) in dichloromethane (20 ml) was added dropwise tert-butyl hypochlorite (0.46 ml) under ice-cooling, and the mixture was stirred under ice-cooling for 50 minutes. The reaction mixture was washed with water and saturated brine, and dried over anhydrous sodium sulfate. The solvent was distilled away under reduced pressure, and the residue was purified by silica gel column chromatography (developing solvent: hexane/ethyl acetate=3/1 v/v) to give the title compound (830 mg, yield 82%).

Step 6) N-[3,5-Dichloro-2-hydroxy-4-(4-methylaminobutylamino)benzoyl]-L-phenylalanine methyl ester dihydrochloride (I)

To a solution of the compound (280 mg) obtained in the above Step 11) in chloroform (5 ml) was added trifluoroacetic acid (2.5 ml) at room temperature, and the mixture was stirred for 20 minutes. Toluene was added to the reaction mixture and the mixture was concentrated under reduced pressure. A 1M hydrogen chloride-ether solution was added to the residue for crystallization to give the title compound (218 mg, yield 82%, see Table 71).

## Example 137

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The compound of Example 137 was prepared in the same manner as in Example 136 from the corresponding compound (see Table 71).

# Example 138

N-[3,5-Dichloro-2-hydroxy-4-(4-aminobutoxy)benzoyl]-L-phenylalanylaminobenz ne hydrochloride

3,5-Dichloro-2-hydroxy-4-(4-tert-butoxycarbonylaminobutoxy)benzoic acid (347 mg) obtained in the same manner

as in the above Exampl 1, Step 4) and L-phenylalanylaminobenzene hydrochloride (268 mg) were subjected to the same reaction as in the above Example 1, Step 5) and Step 6) to give the title compound (284 mg, yield 58%, see Table 72).

## 5 Examples 139-142

The compounds of Examples 139-142 were prepared in the same manner as in Example 138 from the corresponding compounds (see Tables 72-74).

#### 10 Example 143

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N-[3,5-Dichloro-2-hydroxy-4-(4-methylaminobutoxy)benzoyl]-L-phenylalanylhydroxyamide

Step 5) N-[3,5-Dichloro-2-hydroxy-4-(4-benzyloxycarbonylmethylaminobutoxy)benzoyl]-L-phenylalanyl-O-benzylhy5 droxyamide (I')

3,5-Dichloro-2-hydroxy-4-(4-benzyloxycarbonylaminobutoxy)benzoic acid (237 mg) obtained in the same manner as in the above Example 1, Step 4) and L-phenylalanyl-O-benzylhydroxyamide hydroxhloride (203 mg) were subjected to the same reaction as in the above Example 1, Step 5) to give the title compound (325 mg, yield 59%).

Step 6) N-[3,5-Dichloro-2-hydroxy-4-(4-methylaminobutoxy)benzoyl]-L-phenylalanylhydroxyamide (I)

To a solution of the compound (210 mg) obtained in the above Step 5) in methanol (5 ml) was added palladium hydroxide (42 mg), and the mixture was subjected to catalytic hydrogenation using hydrogen gas under atmospheric pressure. Twelve hours later, the reaction mixture was filtered, and the filtrate was concentrated under reduced pressure. Methanol-ether was added to the residue for crystallization to give the title compound (188 mg, yield 62%, see Table 74).

#### Example 144

N-[4-(4-Aminobutoxy)-3,5-dichloro-2-hydroxybenzoyl]-1-(3-methyl-1,2,4-oxadiazol-5-yl)-2-phenylethylamine hydrochloride

3,5-Dichloro-2-hydroxy-4-(4-tert-butoxycarbonylaminobutoxy)benzoic acid (394 mg) obtained in the same manner as in the above Example 1, Step 4) and 1-(3-methyl-1,2,4-oxadiazol-5-yl)-2-phenylethylamine hydrochloride (240 mg) were subjected to the same reaction as in the above Example 1, Step 5) and Step 6) to give the title compound (299 mg, yield 58%, see Table 75).

## Example 145

N-[4-(4-Aminobutoxy)-3,5-dichloro-2-hydroxybenzoyl]-L-phenylalaninol hydrochloride

3,5-Dichloro-2-hydroxy-4-(4-tert-butoxycarbonylaminobutoxy)benzoic acid (394 mg) obtained in the same manner as in the above Example 1, Step 4) and O-benzyl-L-phenylalaninol (242 mg) were subjected to the same reaction as in the above Example 1, Step 5), Example 99, Step 6) and Example 1, Step 6) to give the title compound (190 mg, yield 42%, see Table 75).

### Example 146

(2S)-3-Phenyl-2-[5-(4-aminobutoxy)-3-hydroxy-2-naphthoylamino]propionic acid methyl ester hydrochloride

Step 13) (2S)-3-Phenyl-2-(3,5-dihydroxy-2-naphthoylamino)propionic acid methyl ester (XVII)

A solution of 3,5-dihydroxy-2-naphthoic acid (4.08 g), L-phenylalanine methyl ester hydrochloride (4.74 g), WSC • HCI (4.22 g), HOBT (2.97 g) and N-methylmorpholine (2.41 ml) in DMF (200 ml) was stirred at room temperature for 16 hours. Water was added to the reaction mixture, and the mixture was extracted with ethyl acetate. The organic layer was washed successively with a 10% aqueous citric acid solution, water, a saturated aqueous sodium hydrogencarbonate solution, water and saturated brine, dried over anhydrous sodium sulfate, and concentrated under reduced

pressure. The residue was purified by silica gel column chromatography (developing solvent: hexane/ethyl acetate=1/1 v/v) to giv the title compound (4.42 g, yield 61%).

Step 14) (2S)-3-Phenyl-2-[5-(4-tert-butoxycarbonylaminobutoxy)-3-hydroxy-2-naphthoylamino]propionic acid methyl ester (I')

To a solution of the compound (1.83 g) obtained in the above Step 13), triphenylphosphine (1.31 g) and 4-tert-butoxycarbonylaminobutyl alcohol (473 mg) in THF (25 ml) was added dropwise disopropyl azodicarboxylate (0.98 ml) at room temperature. After stirring at room temperature for 16 hours, the reaction mixture was concentrated under reduced pressure, and the residue was purified by silica gel column chromatography (developing solvent: hexane/ethyl acetate=2/1 v/v) to give the title compound (375 mg, yield 30%).

Step 6) (2S)-3-Phenyl-2-[5-(4-aminobutoxy)-3-hydroxy-2-naphthoylamino]propionic acid methyl ester hydrochloride (I)

To a solution of the compound (375 mg) obtained in the above Step 14) in THF (5 ml) was added a 4N hydrogen chloride-dioxane solution (5 ml), and the mixture was stirred at room temperature for 3 hours. The mixture was concentrated under reduced pressure. Ether was added to the residue for crystallization to give the title compound (187 mg, yield 57%, see Table 76).

## 20 Example 147

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N-[4-[4-(4-Methylaminobutoxy)phenyl]benzoyl]-L-phenylalanine ethyl ester hydrochloride

Step 13) 4-(4-Hydroxyphenyl)benzoyl-L-phenylalanine ethyl ester (XVII)

To a solution of 4-(4-hydroxyphenyl)benzoic acid (3.0 g) and L-phenylalanine ethyl ester hydrochloride (3.38 g) in DMF (30 ml) were added WSC • HCI (2.7 g), HOBT (1.89 g) and triethylamine (2 ml), and the mixture was stirred at room temperature for 14 hours. Water was added to the reaction mixture and the mixture was extracted with ethyl acetate. The organic layer was washed successively with a 10% aqueous citric acid solution, water, a saturated aqueous sodium hydrogen-carbonate solution, water and saturated brine, dried over anhydrous sodium sulfate, and concentrated under reduced pressure to give a crude product of the title compound.

Step 15) N-[4-[4-(4-tert-Butoxycarbonylmethylaminobutoxy)phenyl]benzoyl]-L-phenylalanine ethyl ester (I')

To a solution of the crude product obtained in the above Step 13) in DMF (30 ml) were added 4-(tert-butoxycarbo-nylmethylamino)butyl bromide (4.46 g) and potassium carbonate (4.65 g), and the mixture was stirred at room temperature for 14 hours. Ethyl acetate was added to the reaction mixture. The mixture was washed successively with water, a 10% aqueous citric acid solution and saturated brine, and dried over anhydrous sodium sulfate. The organic layer was concentrated under reduced pressure, and the residue was purified by silica gel column chromatography (developing solvent: hexane/ethyl acetate=3/1 v/v) to give the title compound (967 mg, yield 10%).

Step 6) N-[4-[4-(4-Methylaminobutoxy)phenyl]benzoyl]-L-phenylalanine ethyl ester hydrochloride (I)

To a solution of the compound (140 mg) obtained in the above Step 14) in THF (2 ml) was added a 4N hydrogen chloride-dioxane solution (2 ml). The mixture was stirred at room temperature for 4 hours, and concentrated under reduced pressure. Ether was added to the residue for crystallization to give the title compound (71 mg, yield 58%, see Table 76).

## Example 148

(2S)-3-Phenyl-2-[4-[5-(4-methylaminobutyl)-1,2,4-oxadiazol-3-yl]-2-hydroxybenzoylamino]propionic acid ethyl ester hydrochloride

Step 9) Methyl 2-hydroxybenzoate-4-carboxamide O-(4-tert-butoxycarbonylmethylaminovaleryl) oxime (XV)

A solution of 4-tert-butoxycarbonylmethylaminovaleric acid (255 mg), methyl 2-hydroxybenzoate-4-carboxamide xime (210 mg), WSC • HCI (211 mg) and 4-dimethylaminopyridine (DMAP, 135 mg) in dichloromethane (5 ml) was stirred at room temperature for 16 hours. Water was added to the reaction mixture and the mixture was extracted with

ethyl acetate. The organic layer was washed successively with a 10% aqueous citric acid solution, water, a saturated aqueous sodium hydrogencarbonate solution, water and saturated brine, dried over anhydrous sodium sulfate, and concentrated under reduced pressur. The residue was purified by silica gel column chromatography (developing solvent; hexane/ethyl acetate=1/1 v/v) to giv the title compound (229 mg, yield 54%).

Step 10) 2-Hydroxy-4-[5-(4-tert-butoxycarbonylmethylaminobutyl)-1,2,4-oxadiazol-3-yl]benzoic acid methyl ester (VI')

A solution of the compound (224 mg) obtained in the above Step 9) in toluene (20 ml) was refluxed under heating for 16 hours. The reaction mixture was allowed to cool, and concentrated under reduced pressure. The residue was purified by silica gel column chromatography (developing solvent: hexane/ethyl acetate=3/1 v/v) to give the title compound (148 mg, yield 69%).

Step 4) 2-Hydroxy-4-[5-(4-tert-butoxycarbonylmethylaminobutyl)-1,2,4-oxadiazol-3-yl]benzoic acid (VII)

To a solution of the compound (146 mg) obtained in the above Step 10) in ethanol (10 ml) was added a 1M lithium hydroxide solution (5 ml), and the mixture was refluxed under heating for 2 hours. The reaction mixture was concentrated under reduced pressure, and a 10% aqueous citric acid solution was added to the residue, which was followed by extraction with ethyl acetate. The organic layer was washed with water, and dried over anhydrous sodium sulfate. The organic layer was concentrated under reduced pressure to give the title compound (140 mg, yield 99%).

Step 5) (2S)-3-Phenyl-2-[4-[5-(4-tert-butoxycarbonylmethylaminobutyl)-1,2,4-oxadiazol-3-yl]-2-hydroxyben-zoylamino]propionic acid ethyl ester (I')

A solution of the compound (140 mg) obtained in the above Step 4), L-phenylalanine ethyl ester hydrochloride (92 mg), WSC • HCI (77 mg), HOBT (54 mg) and triethylamine (0.056 ml) in DMF (1.5 ml) was stirred at room temperature for 15 hours. Water was added to the reaction mixture and the mixture was extracted with ethyl acetate. The organic layer was washed successively with a 10% aqueous citric acid solution, water, a saturated aqueous sodium hydrogencarbonate solution, water and saturated brine, dried over anhydrous sodium sulfate, and concentrated under reduced pressure. The residue was purified by silica gel column chromatography (developing solvent: hexane/ethyl acetate=2/1 v/v) to give the title compound (174 mg, yield 85%).

Step 6) (2S) -3-Phenyl-2-[4-[5-(4-methylaminobutyl)-1,2,4-oxadiazol-3-yl]-2-hydroxybenzoylamino]propionic acid ethyl ester hydrochloride (I)

To a solution of the compound (172 mg) obtained in the above Step 5) in THF (2 ml) was added a 4N hydrogen chloride-dioxane solution (2 ml), and the mixture was stirred at room temperature for 3 hours. The reaction mixture was concentrated under reduced pressure, and ether was added to the residue for crystallization to give the title compound (133 mg, yield 87%, see Table 77).

#### 40 Examples 149-151

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The compounds of Examples 149-151 were prepared in the same manner as in Example 148 from the corresponding compounds (see Tables 77-78).

#### 45 Example 152

(2S)-2-[2-(3-Methylaminopropylsulfanyl)benzoxazole-5-carbonylamino]-3-phenylpropionic acid ethyl ester hydrochloride

50 Step 2) 2-(3-tert-Butoxycarbonylmethylaminopropylsulfanyl)-5-ethoxycarbonylbenzoxazole (VI)

To a solution of 5-ethoxycarbonyl-2-mercaptobenzoxazole (670 mg) in DMF was added 60% sodium hydride (126 mg) in oil under ice-cooling, and the mixture was stirred for 30 minutes. A solution of 3-tert-butoxycarbonylmethylaminopropyl chloride (623 mg) in DMF was added to the reaction mixture, and the mixture was stirred with heating at 60°C for 18 hours. Ethyl acetate was added to the reaction mixture, and the organic layer was washed with water and dried over anhydrous sodium sulfate. The organic layer was concentrated under reduced pressure, and the residue was purified by silica gel column chromatography (developing solvent: hexane/ethyl acetate=4/1 v/v) to give the title compound (594 mg, yield 50%).

Step 4) 2-(3-tert-Butoxycarbonylmethylaminopropylsulfanyl)-5-carboxybenzoxazole (VII)

To a mixed solution of the compound (562 mg) obtained in the above Step 2) in ethanol (2 ml)-THF (2 ml) was added a 1M lithium hydroxide solution, and the mixture was stirred with heating at 60°C for 1 hour. The reaction mixture was concentrated under reduced pressure, and ethyl acetate and a 10% aqueous citric acid solution were added. The organic layer was dried over anhydrous sodium sulfate, and concentrated under reduced pressure to give the title compound (465 mg, yield 98%).

Step 5) (2S)-2-[2-(3-tert-Butoxycarbonylmethylaminopropylsulfanyl)benzoxazole-5-carbonylamino]-3-phenylpropionic acid ethyl ester (I')

A solution of the compound (465 mg) obtained in the above Step 4), L-phenylalanine ethyl ester hydrochloride (302 mg), WSC • HCl (250 mg), HOBT (176 mg) and triethylamine (0.18 ml) in DMF was stirred at room temperature for 14 hours. Water was added to the reaction mixture and the mixture was extracted with ethyl acetate. The organic layer was washed successively with a 10% aqueous citric acid solution, water, a saturated aqueous sodium hydrogencarbonate solution, water and saturated brine, dried over anhydrous sodium sulfate, and concentrated under reduced pressure. The residue was purified by silica gel column chromatography (developing solvent: hexane/ethyl acetate=4/1 v/v) to give the title compound (240 mg, yield 40%).

 Step 6) (2S)-2-[2-(3-Methylaminopropylsulfanyl)benzoxazole-5-carbonylamino]-3-phenylpropionic acid ethyl ester hydrochloride (I)

To a solution of the compound (231 mg) obtained in the above Step 5) in THF (5 ml) was added a 4N hydrogen chloride-dioxane solution (5 ml), and the mixture was stirred at room temperature for 4 hours. The reaction mixture was concentrated under reduced pressure, and ether was added for crystallization to give the title compound (136 mg, yield 67%, see Table 79).

## Examples 153-154

30 The compounds of Examples 153-154 were prepared in the same manner as in Example 152 from the corresponding compounds (see Tables 79-80).

## Example 155

5 N-[3,5-Dichloro-2-hydroxy-4-(3-piperazinylpropionyloxy)benzoyl]-L-phenylalanine ethyl ester dihydrochloride

Step 18) N-[3,5-Dichloro-2-hydroxy-4-[3-(4-tert-butoxycarbonylpiperazinyl)propionyloxy]benzoyl]-L-phenylalanine ethyl ester (I')

To a solution of N-(3,5-dichloro-2,4-dihydroxybenzoyl)-L-phenylalanine ethyl ester (398 mg) obtained in the same manner as in the above Example 1', Step 13), 3- (4-tert-butoxycarbonylpiperazinyl)propionic acid (258 mg) and 4-dimethylaminopyridine (147 mg) in DMF (4 ml) was added WSC • HCl (230 mg) under ice-cooling, and the mixture was stirred at room temperature for 2 hours. Ethyl acetate (40 ml) was added to the reaction mixture, and the mixture was washed successively with water, a saturated aqueous sodium hydrogencarbonate solution, water and saturated brine. The reaction mixture was dried over anhydrous sodium sulfate, and the solvent was distilled away under reduced pressure. The residue was purified by silica gel column chromatography (developing solvent: ethyl acetate/hexane=1/1 v/v) to give the title compound (258 mg, yield 40%).

Step 6) N-[3,5-Dichloro-2-hydroxy-4-(3-piperazinylpropionyloxy)benzoyl]-L-phenylalanine ethyl ester dihydrochloride (I)

To a solution of the compound (258 mg) obtained in the above Step 18) in dichloromethane (2 ml) was added trifluoroacetic acid (2 ml), and the mixture was stirred at room temperature for 10 minutes. The solvent was distilled away under reduced pressure, and 1M hydrogen chloride-ether (3 ml) was added for crystallization to give the title compound (173 mg, yield 70%, see Table 81).

#### **Examples 156-158**

50

The compounds of Examples 156-158 were prepared in the same manner as in Example 155 from the correspond-

ing compounds (see Tables 81-82).

The structures and physical properties of the compounds of the above Examples are shown in the following Tables 2-82.

In the Tables, Me, Et, Ph, Bn and Ac mean methyl, ethyl, phenyl, benzyl and acetyl, respectively.

	Elemental analysis (%)	C <sub>22</sub> H <sub>2</sub> ¢Cl <sub>2</sub> N <sub>2</sub> O <sub>6</sub> ·HCl Calculated C, 52. 24 H, 5. 38 N, 5. 53 Found C, 52. 05 H, 5. 37 N, 5. 51	C16H3, C12N3O5.2HC1 Calculated C, 50.27 H, 5.57 N, 7.03 Pound C, 50.19 H, 5.74 N, 6.93
	FAB-MS	469 (free base, MH+)	524 (free base, MH*)
	IR (cm <sup>-1</sup> )	KBr 3422 2953 1742 1637 1458 1219	KBr 3406 2357 2372 1736 1642 1458
Table 2	'H-NMR & (ppm), 300MHz	DMSO-d <sub>4</sub> 1. 82(4H, bs) 2. 56(3H, t, J=5, 4Hz) 2. 96(2H, bs) 3. 04-3. 28(2H, m) 3. 66(3H, s) 4. 05(2H, bs) 4. 72-4. 82(1H, m) 7. 18-7. 30(5H, m) 8. 17(1H, s) 8. 48(2H, bs) 9. 44(1H, bs) 13. 35(1H, s)	DMSO-d <sub>4</sub> 1. 14(3H, t, J=6. 0Hz) 2. 81(3H, s) 3. 0-3. 60(10H, m) 4. 11(2H, q, J=6. 0Hz) 4. 34(2H, brs) 4. 68-4. 78(1H, m) 7. 19-7. 29(5H, m) 8. 22(1H, s) 9. 46(1H, d, J=7. 0Hz) 13. 40(1H, brs)
	Compound	C1 OH Ph MeN-(CH <sub>2</sub> ), -0 — CONH — COOMe -HC1	$\begin{array}{c} \text{Cl} & \text{OH} \\ \text{Me-N} & \text{N-(CH_2)_2-0} \\ \text{Cl} \\ \text{C2HC1} \end{array}$
	Ex. No.	<u>.</u>	23

	Blemental analysis (%)		
. 15	PAB-MS	372 (free base, MH*)	386 (free base, MH+)
20	[R (cm <sup>-1</sup> )	KBr 3383 1739 1632 1607 1534 1498	KBr 3378 1630 1604 1534 1498
<sup>25</sup> ४. व प	H-NMR & (ppm), 300MHz	74 (2H, m) 72 (2H, m) 73 (2H, m) 8) 74 (1H, m) 74 (1H, m) 74 (5H, m) 75 (74 (5H, m) 76 (74 (5H, m) 77 (74 (5H, m) 77 (74 (5H, m) 78 (74 (5H, m) 79 (74 (5H, m) 79 (74 (5H, m) 70 (74 (5H, m) 71 (74 (5H, m) 71 (74 (5H, m) 72 (74 (5H, m) 74 (5H, m) 74 (5H, m) 74 (5H, m) 75 (74 (5H, m) 76 (74 (5H, m) 77	0-ds 1-1. 73(4H, m) 1-2. 86(2H, m) 1-3. 18(2H, m) 1-4. 11(2H, m) 1-4. 89(1H, m) 1-7. 34(5H, m) 1-7. 34(5H, m) 1-7. 34(5H, m) 1-8. 04(3H, brs) 1(1H, d, J=7. 0Hz) 4(1H, s)
30	'H-NMR &	DMSO-da 1. 86-2. C 1. 82-2. S 3. 02-3. I 3. 64(3H, 4. 06-4. I 4. 72-4. 8 6. 42-6. 5 7. 16-7. 3 7. 65(1H, 7. 96(3H, 8. 21(1H, 10. 24(1H,	DMS0-de 1. 46-1. 7 2. 63-2. 8 3. 00-3. 1 3. 92-4. 1 4. 78-4. 8 6. 43-6. 5 7. 10-7. 3 7. 82-8. 0 8. 22(1H.
35		Ph COOMe	Ph COOMe
40	Compound	OH CONF	HO CONF
45		H2N-(CH2)3-0-	H <sub>2</sub> N-(CH <sub>2</sub> ),-0-
50	RX.	د ج	4 rgH .

5	Elemental analysis (%)		
10	FAB-MS	400 (free base. MH+)	414 (free base, MH*)
	IR (cm <sup>-1</sup> )	KBr 1630 1604 1534 1201	KBr 3378 1630 1605 1534 11498 1181
20	1z		Hz)
Table 4	'H-NMR & (ppm), 300MHz	DMSO-d <sub>6</sub> 1. 18-1. 42(2H, m) 1. 43-1. 54(4H, m) 2. 62-2. 78(2H, m) 3. 02-3. 21(2H, m) 3. 67(3H, s) 3. 91-4. 08(2H, m) 4. 82-4. 94(1H, m) 6. 42-6. 52(2H, m) 7. 08-7. 18(2H, m) 7. 08-7. 18(2H, m) 7. 08-7. 36(3H, m) 7. 20-7. 36(3H, m)	DMSO-d <sub>6</sub> 1. 14-1. 34(4H, m) 1. 40-1. 61(4H, m) 2. 66-2. 80(2H, m) 3. 01-3. 16(2H, m) 4. 80-4. 90(1H, m) 6. 44(1H, d. J=2. 2Hz) 7. 10-7. 32(5H, m) 7. 77(1H, d. J=8. 4Hz) 7. 85(3H, brs) 8. 24(1H, d. J=7. 4Hz) 10. 22(1H, brs)
35	Compound	OH COOME	OH COOMe
40 45	Сомр	H₂N-(CH₂) 6-0 —()	H₂N-(CH₂),•-0 —(
50	BX.	D.	့

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5	Elemental analysis (%)	·	·
	FAB-MS	435 (free base, MH+)	449 (free base. MH+)
	IR (cm <sup>-1</sup> )	Neat 2954 1728 1642 1589 1548 1497	Neat 2958 1773 1641 1588 1547 1497
20	MH2		
rable 5	'H-NMR & (ppm), 300MHz	DNSO-d <sub>6</sub> 1. 72-1. 88(4H, m) 2. 48-2. 57(3H, m) 2. 90-3. 01(2H, m) 3. 10-3. 25(2H, m) 3. 66(3H, s) 4. 16(2H, t, J=6Hz) 4. 69-4. 76(1H, m) 6. 78(1H, d, J=9Hz) 7. 16-7. 32(5H, m) 7. 94(1H, d, J=9Hz) 8. 70(2H, brs) 9. 26(1H, d, J=9Hz) 13. 35(1H, s)	DMSO-d <sub>6</sub> 1. 14(3H, t, J=6Hz) 1. 72-1. 88(4H, m) 2. 49-2. 55(3H, m) 2. 90-3. 02(2H, m) 3. 10-3. 24(2H, m) 4. 11(2H, q, J=6Hz) 4. 17(2H, t, J=6Hz) 4. 65-4. 73(1H, m) 6. 79(1H, d, J=9Hz) 7. 17-7. 32(5H, m) 7. 95(1H, d, J=6Hz) 8. 09(2H, brs) 9. 23(1H, d, J=6Hz) 13. 37(1H, s)
30		-Ph -C00Me	Ph COOBt
35	Compound	OH CONH	OH CONH
40	Com		
<b>45</b>		CI MeN-(CH <sub>2</sub> ) 4-0 - H	MeN-(CH <sub>2</sub> ),-0 H •HCl
	Š.	7	∞

	8		
5	Elemental analysis	·	
10		92.	့် (
45	FAB-MS	435 (free base, MH*)	449 (free base, MH*)
15			
	IR (cm <sup>-1</sup> )		KBr 2950 2783 1745 1637 1544 1465 1369 1264
20	MHz		^
7able 6	'H-NMR & (ppm), 300MHz	DMSO-d <sub>6</sub> 1. 70-1. 86(4H, m) 2. 53(3H, s) 2. 92-3. 02(2H, m) 3. 05-3. 23(2H, m) 3. 65(3H, s) 4. 07-4. 17(2H, m) 4. 68-4. 78(1H, m) 6. 65(1H, s) 7. 20-7. 31(5H, m) 8. 02(1H, s) 8. 99(1H, d, J=7. 0Hz) 12. 49(1H, brs)	DMSO-de 1. 78-1. 84(4H, m) 2. 09(3H, s) 2. 09(3H, s) 2. 53(3H, bs) 2. 95(2H, bs) 3. 08-3. 24(2H, m) 3. 66(3H, s) 3. 88-3. 94(2H, m) 4. 68-4. 82(1H, m) 7. 18-7. 32(5H, m) 7. 18-7. 32(5H, m) 8. 03(1H, s) 8. 78(2H, bs) 9. 29(1H, d. J=7. 7Hz) 12. 92(1H, s)
30		DMe	Ph COOMe
35	Compound	OH CONH COO	OH CONH COO
40	Comi		
45		MeN-(CH <sub>2</sub> ),-0 — H •HC1	MeN-(CH <sub>2</sub> ),-0—(HC1
	Bx.	တ	. 10

5		Elemental analysis (%)	C2.0H22C12N2O6.HC1 Calculated C, 50. 28 H, 4. 85 N, 5. 86 Pound C, 50. 19 H, 4. 69 N, 5. 74	C <sub>2.1</sub> H <sub>24</sub> Cl <sub>2</sub> N <sub>2</sub> O <sub>6</sub> :HCl Calculated C, 51. 29 H, 5. 12 N, 5. 70 C, 50. 78 H, 5. 17 N, 5. 58
15		FAB-MS	441 (free base, MH+)	455 (free base, NH*)
20		IR (cm <sup>-1</sup> )	KBr 3422 2952 2730 1744 1942 1585 1545 1353 1221	KBr 2953 1641 1585 1585 1542 1355 1221
25	Table 7	'H-NMR & (ppm), 300MHz	DMSO-d <sub>4</sub> 2. 65-2. 69(3H, m) 3. 09-3. 22(2H, m) 3. 36-3. 42(2H, m) 3. 64-3. 67(3H, m) 4. 28(2H, t, J=6. 0Hz) 4. 74-4. 81(1H, m) 7. 19-7. 29(5H, m) 8. 23(1H, s) 8. 95-9. 03(2H, m) 9. 53(1H, s) 13. 38(1H, s)	DMSO-d <sub>6</sub> 1. 65-1. 95(4H, m) 2. 77-2. 94(2H, m) 3. 15(1H, dd. 3. 24(1H, dd.
35			NH COOMe	NH COOMe
40		Compound	C1 OH	10 CI OH
45			MeN-(CH <sub>2</sub> ) <sub>2</sub> - H	H <sub>2</sub> N-(CH <sub>2</sub> ),
50		RX.	=	12

5	Elemental analysis (%)	•	
15	FAB-MS	469 (MH+)	483 (free base, MH*)
20	IR (cm <sup>-1</sup> )	KBr 3423 2951 1743 1618 1571 1541 1434 1065	
	Table 8 H-NMR & (ppm), 300MHz	(4H, m) ) (4H, m) ) , J=6.0Hz) (1H, m) (6H, m) rds)	(4H, m) (4H, m) (5H, m) (5H, m) (5H, m)
30	T H-NMR S (	DMSO-de 1. 63-1. 95(4H, m) 2. 55(3H, s) 2. 92-3. 07(4H, m) 3. 57(3H, s) 3. 88(2H, t. J=6. 0Hz) 4. 65-4. 71(1H, m) 7. 18-7. 30(6H, m) 7. 50(1H, s) 8. 41(1H, brds) 12. 25-12. 27(1H, m)	DMSO-de 1. 84 (4H, s) 2. 54 (3H, s) 2. 90-3. 25 (4H, m) 3. 58 (3H, s) 3. 68 (3H, s) 4. 02 (2H, m) 4. 76 (1H, m) 7. 20-7. 32 (5H, m) 7. 41 (1H, m) 8. 75 (1H, d, J=9Hz)
35		COOMe	Ph COOMe
40	Compound	OH	OMe CONH
. 45		MeN-(CH <sub>2</sub> ),4-0 — C1.	C1. MeN-(CH <sub>2</sub> ),-0 — H C1.
	Bx. No.	Met 13	Mel 14
	L		

5

Elemental analysis (%)	C <sub>22</sub> H <sub>2s</sub> Cl <sub>2</sub> N <sub>2</sub> O <sub>6</sub> ·HCl Calculated C, 50.64 H, 5.22 N, 5.37 Found C, 49.62 H, 5.29 N, 5.46
PAB-MS	485 (free base, MH+)
IR (cm-1)	KBr 1640 1586 1515 1458 1221
'H-NMR & (ppm), 300MHz	DMSO-ds 1. 79-1. 89(4H, m) 2. 55(2H, t, J=6. 0Hz) 2. 85-3.00(2H, m) 3. 06(1H, dd, J=15. 6, 8. 4Hz) 3. 57(3H, s) 3. 57(3H, s) 4. 01-4. 11(1H, m) 4. 50-4. 53(2H, m) 4. 59-4. 71(1H, m) 6. 66(2H, d, J=6. 0Hz) 7. 06(2H, d, J=6. 0Hz) 8. 20(1H, brs) 8. 65(2H, brs) 9. 26(1H, s) 13. 37(1H, s)
Compound	MeN-(CH <sub>2</sub> ),-0-0H
RX.	15

Table 9

5		Elemental analysis (%)	C11418C11N2O5-HC1 Calculated C, 53. 14 H, 5. 62 N, 5. 39 Found C, 52. 54 H, 5. 50 N, 5. 40	·
15		FAB-MS	483 (free base. MH⁺)	483 (MH*)
20		IR (cm <sup>-1</sup> )	KBr 2954 1747 1641 1584 1542 1458 1354 1219	Neat 2952 2360 1743 1633 1437
25 30	Table 10	¹H-NMR & (ppm), 300MHz	DMSO-d <sub>8</sub> 1, 20(3H, t, J=6, 0Hz) 1, 82-1, 88(4H, m) 2, 92-2, 95(4H, m) 3, 09-3, 33(2H, m) 3, 66(3H, s) 4, 03-4, 07(2H, m) 4, 71-4, 79(1H, m) 7, 19-7, 29(5H, m) 8, 20(1H, s) 8, 58-8, 76(2H, m) 9, 48(1H, d, J=6, 0Hz) 13, 35(1H, s)	CDC13 1. 80-1. 91 (2H, m) 1. 95-2. 07 (2H, m) 2. 59 (6H, s) 2. 96-3. 13 (2H, m) 3. 18-3. 32 (2H, m) 3. 73 (3H, s) 3. 96-4. 07 (2H, m) 5. 15 (1H, q, J=6Hz) 7. 10-7. 30 (5H, m) 7. 85 (1H, s) 9. 97 (1H, brs)
35		id .	ONH COOMe	CONH COOMe
40		Compound	CI	10 0 10
45			C1. BtN-(CH <sub>2</sub> ),-0 — H	CI N62N-(CH2),4-0 -
50		Š.	91	17

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78	3
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%.S	Compound	¹Н-NMR 6 (ррт), 300MHz	IR (cm <sup>-1</sup> )	FAB-MS	Elemental analysis (%)
82	Me <sub>2</sub> N-(CH <sub>2</sub> ),-0 — CONH — COOMe -HC1	CDC13 1. 90-2. 01(2H, m) 2. 13-2. 25(2H, m) 2. 83(6H, s) 3. 13-3. 30(4H, m) 3. 81(3H, s) 5. 02(1H, q, J=6Hz) 7. 14-7. 43(7H, m) 7. 48(1H, s)	Neat 3241 2955 2671 1743 1640 1584 1461	483 (free base. MH <sup>+</sup> )	C <sub>23</sub> H <sub>28</sub> Cl <sub>2</sub> N <sub>2</sub> O <sub>6</sub> ·HCl Calculated C, 53. 14 H, 5. 62 N, 5. 39 Pound C, 53. 24 H, 5. 63 N, 5. 34
61	C1 OH Ph H <sub>2</sub> N-(CH <sub>2</sub> ), -0 — CONH — C00Bt -HC1	DMSO-ds 1. 14(3H, t, J=6. 0Hz) 1. 70-1. 95(4H, m) 2. 80-2. 95(2H, m) 3. 95-4. 15(4H, m) 4. 60-4. 75(1H, m) 7. 18-7. 40(5H, m) 7. 91(3H, brs) 8. 21(1H, s) 9. 47(1H, d, J=6. 0Hz) 13. 36(1H, s)	KBr 2961 1722 1722 1643 1544 1459 1354	469 (free base, MH+)	C <sub>22</sub> H <sub>2</sub> C <sub>12</sub> N <sub>2</sub> O <sub>3</sub> ·HC <sub>1</sub> Calculated C, 52.34 H, 5.19 N, 5.55 Found C, 51.62 H, 5.41 N, 5.48

5	Elemental analysis (%)	C1141201200-HC1 Calculated C, 53. 14 H, 5. 62 N, 5. 39 Found C, 52. 85 H, 5. 69 N, 5. 24	
. 10	Elemen		
15	FAB-MS	483 (free base, MH+)	483 (MH+)
20	[R (cm <sup>-1</sup> )	KBr 3422 2959 1736 1627 1447 1333 1182	
rable 12	'H-NMR & (ppm), 300MHz	-d; 3H, t. J=7Hz) 4H, brs) 3. 39(7H, m) 2H, brs) 2H, brs) 6. 7H, m) 6. 7H, m) 6. 3H, brs) 6. 7H, brs) 7. 39(5H, m) 3H, brs) (1H, brs)	CDC1. 1. 16(3H. t. J=8Hz) 1. 50-1. 75(4H, m) 2. 38(3H. s) 2. 30-3. 05(2H, m) 3. 26(2H, dq. J=3. 12Hz) 3. 30-3. 45(2H, m) 4. 00-4. 10(2H, m) 5. 02-5. 10(1H, m) 7. 10-7. 15(2H, m) 7. 20-7. 30(3H, m) 8. 00(1H, s) 10. 76(1H, brs)
30	MN-H1	DMSO-(38) 1. 20(38) 1. 81(48) 1. 81(48) 2. 55-34 3. 96(28) 6. 08(0,6) 6. 08(0,6) 6. 94-7,7 7. 91(38)	
35		N COOB!	NH COOE
40	Compound	10 10 10 10	10 C1 C1
45		C1 H2N-(CH2),4-0 — ·HC1	C1. MeN-(CH <sub>2</sub> ),-0— C1'
50	S.S.	50	22

5	Blemental analysis (%)	C <sub>23</sub> H <sub>2</sub> gCl <sub>2</sub> N <sub>2</sub> O <sub>6</sub> ·HCl Calculated C, 53. 14 H, 5. 62 N, 5. 39 Pound C, 53. 36 H, 5. 71 N, 5. 53	
15	FAB-MS	483 (free base, MH+)	497 (MH+)
20	IR (cm <sup>-1</sup> )	KBr 1740 1584 1459 1352 1216	Neat 2956 1738 1634 1574 1538 1440
72 Table 13	'H-NAIR & (ppm), 300MHz	DMSO-de 1. 14(3H, t, J=6. 0Hz) 1. 77-1. 91(4H, m) 2. 554(3H, t, J=6. 0Hz) 2. 89-3. 00(2H, m) 3. 13(1H, dd, J=9. 0, 15. 0Hz) 3. 22(1H, dd, J=6. 15. 0Hz) 4. 00-4. 11(2H, m) 4. 08(2H, q, J=6. 0Hz) 4. 68-4. 79(1H, m) 7. 18-7. 32(5H, m) 8. 21(1H, s) 8. 21(1H, s) 9. 48(1H, d, J=6. 9Hz) 13. 36(1H, s)	CDC1s 1. 27(3H, t, J=7. 5Hz) 1. 82-2. 04(4H, m) 2. 55(6H, s) 2. 95-3. 11(2H, m) 3. 25(2H, d, J=4Hz) 3. 93-4. 04(2H, m) 4. 12-4. 22(2H, m) 5. 11-5. 18(1H, m) 7. 13-7. 30(5H, m) 7. 90(1H, s) 10. 31(1H, brs)
35		Ph COOBt	NA Ph
40	Compound	10 10 10	15 CI
45		MeN-(CH <sub>2</sub> ),-0· H (C	Me <sub>2</sub> N-(CH <sub>2</sub> ),-0
50	Bx.	83	g

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/sis	101	101
Elemental analysis (%)	C <sub>22</sub> H <sub>2</sub> Cl <sub>2</sub> N <sub>2</sub> O <sub>5</sub> ·HCl Calculated C, 53. 99 H, 5. 85 N, 5. 25 Round C, 54. 11 H, 5. 86 N, 5. 27	C24H3,C12N2O6.HC1 Calculated C,53.99 H, 5.85 N, 5.25 Found C,52.75 H, 5.59 N, 4.72
nental (%	12,012 2012,012 53.99 54.11 54.11 5.27	4. 72
Elen	Call Call N.H.C. E	22 24 25 25 25 25 25 25 25 25 25 25 25 25 25
FAB-MS	base. MH+)	497 (free base. MH*)
FAB	497 (free base, MH*)	(free
[R (cm <sup>-1</sup> )	Neat 2956 1738 1639 1583 1461	KBr 1641 1585 1458 1219
	-8111	
300MHz	5HZ)	0Hz)
(ppm),	1, J=7, 5Hz 35(4H, m) 2, s) 24(4H, m) 14(4H, m) 75(1H, m) 75(1H, m) 75(1H, m) 75(1H, m) 75(1H, m) 75(1H, m)	25 (6.1) 1-6 (7.1) 1-6 (7.1) 1-6 (7.1) 1-6 (7.1) 1-6 (7.1) 1-7 (7.1) 1-7 (7.1) 1-7 (7.1) 1-7 (7.1) 1-6 (7.
H-NMR & (ppm), 300MHz	DMSO-ds 1. 13(3H, t. J=7.5H 1. 76-1. 95(4H, m) 2. 74(6H, s) 3. 06-3. 24(4H, m) 4. 04-4. 14(4H, m) 7. 18-7. 29(5H, m) 7. 18-7. 29(5H, m) 8. 21(1H, s) 9. 54(1H, brs)	DMSO-de 1. 13(3H, t, J=6. 0Hz) 1. 30-1. 62(6H, m) 1. 65-1. 80(2H, m) 2. 80-2. 88(2H, m) 3. 03-3. 27(2H, m) 4. 60-4. 78(1H, m) 7. 10-7. 40(5H, m) 7. 78(3H, brs) 8. 19(1H, s) 9. 44(1H, d, J=6. 0Hz) 13. 35(1H, s)
_		
	- C00Et	1 COORT
	HNOO.	
Compound	HO HO	HO L
Com		
	H <sub>2</sub> ), -0	0-
	MezN-(CHz)	H2N-(CH2)
S. S.	24	52
<u> </u>	<u> </u>	<u> </u>

		<del></del>	
5	Elemental analysis (%)	Ca 6H3 Cl 2N2 O6 · HCl Calculated C, 54.80 H, 6.07 N, 5.11 Found C, 53.81 H, 6.10 N, 4.96	C23H26C12N2O6.HC1 Ca1culated C, 53.14 H, 5.62 N, 5.39 Found C, 51.51 H, 5.41 N, 4.99
15	FAB-MS	511 (free base, MH*)	483 (free base, MH+)
20	IR (cm <sup>-1</sup> )	KBr 3420 2936 1719 1641 1585 1543 1458 1219	KBr 3420 2981 1717 1641 1585 1458
Table 15	'H-NMR & (ppm), 300MHz	DMSO-d <sub>4</sub> 1. 13(3H, t, J=7. 1Hz) 1. 33-1. 36(4H, m) 1. 47-1. 58(4H, m) 1. 47-1. 58(4H, m) 2. 72-2. 82(2H, m) 3. 08-3. 26(2H, m) 4. 03(2H, t, J=6. 4Hz) 4. 11(2H, q, J=7. 1Hz) 4. 66-4. 73(1H, m) 7. 18-7. 29(5H, m) 7. 18-7. 29(5H, m) 7. 73-7. 84(3H, m) 8. 19(1H, d, J=7. 1Hz) 9. 456(1H, d, J=7. 1Hz) 13. 36(1H, S)	DMSO-de 1. 08-1. 10(3H, d, J=6. 0Hz) 1. 17-1. 19(3H, d, J=6. 0Hz) 1. 82(4H, brs) 2. 88(2H, brs) 3. 10-3. 30(2H, m) 4. 04(2H, brs) 4. 60-4. 90(2H, m) 7. 21-7. 30(5H, m) 7. 21-7. 30(5H, m) 7. 89(3H, brs) 8. 19(1H, s) 9. 50(1H, brs) 13. 38(1H, brs)
30	N-H <sub>1</sub>	DMSO-de 1. 13(3H, 1 1. 33-1. 36 1. 47-1. 58 1. 77-2. 82 3. 08-3. 26 4. 03(2H, 1 4. 66-4. 73 7. 18-7. 29 7. 73-7. 84 8. 19(1H, 5, 9, 9, 45(1H, 6, 1)	DMSO- 1. 08-1. 1. 17-1. 1. 82(4) 2. 88(2) 3. 10-3. 4. 60-4. 7. 21-7. 7. 21-7. 7. 89(3) 8. 19(1) 9. 50(1)
35		Ph COOB!	Ph 600
40	Compound	OH CONH	HO COM
45		C1 H2N-(CH2), -0 —< C1	C1 H2N-(CH2),-0— C1
50			
	BX.	82	27

5	Elemental analysis (%)	C24H30C12N2O8·HC1 Calculated C, 55.99 H, 5.85 N, 5.25 Found C, 53.22 H, 5.94 N, 5.21	C2sH3zCl2N2Os-HCl Calculated C. 54.80 H. 6.07 N. 5.11 Pound C. 54.59 H. 6.06 N. 4.98
15	FAB-MS	497 (free base, MH+)	511 (free base. MH <sup>+</sup> )
20	IR (cm <sup>-1</sup> )	KBr 3385 2962 1721 1642 1585 1542 1458 1355 1218	KBr 3360 2361 1740 1640 1584 160
25 Table 16	'H-NMR & (ppm), 300MHz	DMSO-d <sub>6</sub> . 0. 82(6H, d. J=6. 0Hz) 1. 74-1. 90(4H, m) 2. 80-2. 95(2H, m) 3. 10-3. 28(3H, m) 3. 86(2H, d. J=6. 0Hz) 4. 06-4. 10(2H, m) 4. 70-4. 78(1H, m) 7. 16-7. 32(5H, m) 7. 85(3H, brs) 8. 19(1H, s) 9. 45(1H, d. J=7. 0Hz) 13. 37(1H, s)	DMSO-d <sub>6</sub> 0. 82(6H, d. J=6. 7Hz) 1. 75-1. 90(5H, m) 2. 54(3H, s) 2. 90-3. 30(4H, m) 3. 85(2H, d. J=7. 0Hz) 4. 00-4. 10(2H, m) 4. 60-4. 70(1H, m) 7. 15-7. 32(5H, m) 8. 19(1H, s) 8. 67(2H, brs) 9. 50(1H, brs) 13. 38(1H, s)
35	pu	ONH COO	ONH COO
40	Compound	13	13 0
45		C1, H2N-(CH2),-0 — C1'	C1. MeN-(CH <sub>2</sub> ), -0 — H .HC1
50	Ex.	82	29

5	Elemental analysis (%)	Ca4H3,Cl2N2O5.HCl Calculated C, 53, 99 H, 5, 85 N, 5, 25 Found C, 53, 83 H, 6, 14 N, 5, 07	C2.445.C12NaOs.+C1 Calculated C, 57, 00 H, 6. 66 N, 4. 75 Found C, 56. 96 H, 6. 83 N, 4. 53
15	FAB-MS	497 (free base, MH+)	553 (free base, MH*)
20	IR (cm <sup>-1</sup> )	KBr 2977 1640 1586 1386 1153	KBr 3423 2957 2856 1741 1638 1584 1541 1461 1461 1259 1226
Table 17	1H-NMR & (ppm), 300MHz	DNSO-d <sub>8</sub> 1. 35(9H, s) 1. 70-1. 94(4H, m) 2. 77-3. 01(2H, m) 3. 05-3. 18(2H, m) 4. 00-4. 10(2H, m) 4. 52-4. 68(1H, m) 7. 15-7. 34(5H, m) 7. 80-8. 03(3H, brs) 8. 21(1H, s) 9. 40(1H, brs) 13. 44(1H, s)	DMSO-d <sub>4</sub> 0.81(3H, t, J=6.0Hz) 1.12-1.24(8H, m) 1.44-1.54(2H, m) 1.78-1.89(4H, m) 2.53-2.57(3H, m) 2.91-2.98(2H, m) 3.10-3.25(2H, m) 4.05(4H, t, J=6.0Hz) 4.68-4.75(1H, m) 7.16-7.35(5H, m) 8.20(1H, s) 8.70-8.78(2H, m) 9.48(1H, d, J=9.0Hz) 13.40(1H, s)
30		1.1.2.8.4.4.7.7.8.9.9.1.1.3.	
35	I	H- X-	Ph C00 (CH <sub>2</sub> ) 6Me
40	Compound	HO OH	OH
45		C1) -4C1	MeN-(CH <sub>2</sub> ), -0 -1 -1 -1 -1 -1 -1 -1 -1 -1 -1 -1 -1 -1
50 .	Ex. No.	30	31

5		Elemental analysis (%)		
15	·	FAB-MS	497 (free base, MH+)	511 (free base, MH+)
20		(cm <sup>-1</sup> )		Neat 3345 1721 1644 1584 1457
25	Table 18	¹H-NMR & (ppm), 300MHz	DMSO-d <sub>8</sub> 1. 64-1. 87(4H, m) 3. 07-3. 27(4H, m) 3. 67(3H, s) 4. 05(2H, t, J=6Hz) 4. 71-4. 81(1H, m) 6. 90-7. 60(10H, m) 7. 73(1H, t, J=6Hz) 8. 20(1H, s) 9. 50(1H, d, J=9Hz) 13. 35(1H, s)	DMSO-d <sub>4</sub> 1. 14(3H, t, J=7. 5Hz) 1. 65-1. 86(4H, m) 3. 10-3. 25(4H, m) 4. 05(2H, t, J=6Hz) 4. 11(2H, q, J=6Hz) 4. 68-4. 77(1H, m) 6. 88-7. 60(10H, m) 7. 76(1H, t, J=6Hz) 8. 22(1H, s) 9. 49(1H, d, J=9Hz) 13. 36(1H, s)
35	٠		NH COOMe	WH COOEt
40		Compound	10 10 10	13 O-
45			H <sub>2</sub> N-C-N-(CH <sub>2</sub> ),           -  -  -  -  -  -  -  -  -  -	H <sub>2</sub> N-C-N-(CH <sub>2</sub> ),           -  -  -  -  -  -  -  -  -  -
50		Bx. No.	ж 325	£ £

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S.	Compound	'H-NMR & (ppm), 300MHz	IR (cm <sup>-1</sup> )	PAB-MS .	Blemental analysis (%)
£.	MeN-(CH <sub>2</sub> ), -0 — CONH — COOMe H -HC1	DMSO-de 0.80-1.48(5H m) 1.50-1.90(10H m) 2.55(3H, t, J=5.3Hz) 2.96(1H, brs) 3.66(3H, s) 4.04-4.12(2H m) 4.52-4.62(1H, m) 8.29(1H, s) 8.68(2H, brs) 9.31(1H, d, J=6.8Hz) 13.54(1H, s)	KBr 3290 2925 1750 1584 1461 1225	475 (free base, MH+)	C <sub>22</sub> H <sub>32</sub> Cl <sub>2</sub> N <sub>2</sub> O <sub>6</sub> ·HCl Calculated C, 51, 62 H, 6, 50 N, 5, 47 Pound C, 51, 65 H, 6, 20 N, 5, 73
. 89	C1 OH Ph MeN(CH <sub>2</sub> ) <sub>2</sub> O(CH <sub>2</sub> ) <sub>2</sub> -0 C0MH C00Me	DMSO-d <sub>6</sub> 2. 58(3H, s) 2. 98-3. 13(4H, m) 3. 59(3H, s) 3. 74-3. 83(4H, m) 4. 09(2H, t, J=6.0Hz) 4. 66-4. 74(1H, m) 7. 20-7. 28(7H, m) 7. 53-7. 69(1H, m)	KBr 3424 2952 1743 1625 1542 1435 1209 1067	485 (MH⁺)	·

5	Elemental analysis	C12H16C12N2O6.HC1 Calculated C. 50. 64 H. 5. 22 N. 5. 37 Pound C, 50. 64 H. 5. 13 N. 5. 27	
15	PAB-MS	485 (free base, MH <sup>+</sup> )	499 (MH+)
20	[R (cm <sup>-1</sup> )	KBr 2953 2749 1745 1639 1584 1541 1468 1349 1220	·
Table 20	'H-NMR & (ppm), 300MHz	DMSO-d. 2. 54-2. 58(3H, m) 3. 09-3. 22(4H, m) 3. 66(3H, s) 3. 78(2H, t, J=5. 2Hz) 3. 83(2H, t, J=4. 5Hz) 4. 22(2H, t, J=4. 5Hz) 4. 72-4. 80(1H, m) 7. 18-7. 29(5H, m) 8. 20(1H, s) 8. 20(1H, s) 9. 50(1H, d, J=6. 5Hz) 13. 36(1H, dr ds)	CDC1 <sub>3</sub> 2. 65(6H, S) 2. 96-3. 15(2H, m) 3. 19-3. 32(2H, m) 3. 74(3H, S) 3. 74(3H, S) 3. 78-3. 86(4H, m) 4. 18(2H, t, J=6. 0Hz) 5. 09-5. 13(1H, m) 7. 21-7. 28(4H, m) 7. 21-7. 28(4H, m) 7. 6(1H, S) 9. 18(1H, brds)
35		H COOMe	Fh COOMe
40	Compound	)2-0-CON	10 CI CON
45		MeN(CH2)20(CH3)4 H •HCl	MesN(CH2)20(CH2)20
50	Ex.	Net 1	37 Me

	S		
5	Elemental analysis (%)	·	CashasClaNaOs.HCl Calculated C. 51. 55 H. 5. 45 N. 5. 23 Round C. 51. 49 H. 5. 24 N. 5. 24
10			
15	PAB-MS	499 (MH+)	499 (free base, MH+):
20	IR (cm <sup>-1</sup> )	·	KBr 2978 1743 1638 1584 1540 1469 1260 1214
25 10 10 10 10 10 10 10 10 10 10 10 10 10	H-NMR & (ppm), 300MHz	CDC13 1. 25(3H, t, J=7. 1Hz) 2. 53(3H, s) 2. 91-3. 25(2H, m) 3. 21-3. 25(2H, m) 3. 78-3. 85(2H, m) 3. 78-3. 82(2H, m) 4. 10-4. 18(4H, m) 5. 11-5. 17(1H, m) 7. 13-7. 26(6H, m) 8. 00(1H, s) 11. 18(1H, brds)	DMSO-da 2. 56(3H, brds) 2. 56(3H, brds) 3. 10-3. 26(4H, m) 3. 76(2H, t. J=5. 0Hz) 3. 81-3. 85(2H, m) 4. 21-4. 25(2H, m) 4. 69-4. 76(1H, m) 7. 19-7. 30(5H, m) 8. 22(1H, s) 8. 71(2H, m) 9. 55-9. 57(1H, m) 13. 38(1H, brds)
30	=		
35		ONT Ph.	ONH COOB!
40	Compound	13 0-1	10 C1
45		MeN(CH <sub>2</sub> ) <sub>2</sub> 0(CH <sub>2</sub> ) <sub>2</sub> .	MeN(CH2)20(CH2)2- H •HC1
50			
	8. 8. 8.	88	33

`*5*5

56

5	Elemental analysis (%)		
	FAB-MS	499 (free base, MH+)	527 (free base, MH <sup>+</sup> )
15	[R (cm <sup>-1</sup> )		
20	H-NAIR & (ppm), 300AHz	rrs) (4H, m) (2H, m) (2H, m) (2H, m) (2H, m) (1H, m) (5H, m) (5H, m)	J=6. 2Hz) s) J=5. 3Hz) 6H, m) 5H, m)
Table 22	H-NAR & (p	DMSO-de 1. 84(4H, brs) 2. 54(3H, s) 2. 95(2H, brs) 3. 09-3. 40(4H, m) 3. 33(3H, s) 3. 50-3. 60(2H, m) 4. 05(2H, brs) 4. 11(2H, t, J=6Hz) 4. 74-4. 84(1H, m) 7. 20-7. 30(5H, m) 8. 22(1H, s) 8. 74(2H, brs) 9. 50(1H, s)	DMSO-d <sub>e</sub> 1. 03(3H, t, J=6. 2Hz) 1. 83(4H, brs) 2. 53(3H, t, J=5. 3Hz) 2. 80-3. 60(6H, m) 4. 05(2H, m) 4. 20(2H, m) 4. 76(1H, m) 7. 20-7. 40(5H, m) 8. 19(1H, s) 8. 55-8. 85(2H, m) 9. 48(1H, br) 13. 37(1H, s)
30		)(CH2),0H	0(CH2)20Bt
35	puno	Ph COOM	A ESO
40	Compound	C1 OH C1 OH	10 C1 C1
<b>45</b> .	·	MeN-(CH2)4- H +HC1	C1. MeN(CH2)4-0 — H HC1
50	Ex. No.	40	41

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5	Elemental analysis (%)		
10	FAB-MS	555 (free base, MH+)	527 (free base. MH+)
15	IR (cm <sup>-1</sup> )		KBr 3426 2960 1751 1640 1585 1458 1178
20	m), 300MHz	-d <sub>6</sub> 31. 94(4H, m) 33. 02(2H, m) 34. 28(2H, m) 3. 42(2H, m) 3. 42(2H, m) 3. 42(2H, m) 4. 21(2H, m) 7. 35(5H, m) 1H, 3) 1H, 4, J=5. 0Hz) (1H, d, J=5. 0Hz)	(a, t, j=7, 4Hz) (a, t, j=7, 4Hz) (a, t, m) (a, t, m) (a, t, m) (a, t, m) (a, t, m) (a, t, m) (b, m) (c, t, m)
rable 23	'H-NMR & (ppm), 300MHz	DMSO-de 1. 72-1. 94(4H. m. 1. 72-1. 94(4H. m. 1. 72-1. 94(4H. m. 2. 55(3H. t. J=5. m. 2. 20(3H. s.) 3. 20(3H. s.) 3. 20(3H. s.) 3. 30-3. 28(2H. m. 3. 30-3. 40(2H. m. 3. 93-4. 08(2H. m. 4. 09-4. 21(2H. m. 5) 3. 37(1H. s.) 13. 37(1H. s.)	DMSO-ds 1. 20(3H, t, J=7. 4) 1. 80-1.84(4H, m) 2. 82-2.92(2H, m) 3. 14-3.32(2H, m) 4. 00-4.04(2H, m) 4. 12(2H, q, J=7. 4) 4. 72(2H, g, J=7. 4) 4. 72(2H, g, J=7. 4) 7. 16-7. 34(5H, m) 7. 16-7. 34(5H, m) 8. 21(1H, s) 9. 54(1H, d, J=8. 8) 13. 33(1H, s)
s Tabl	•		
35	F	-Ph -C00(CH <sub>2</sub> ) <sub>2</sub> 0(CH <sub>2</sub> ) <sub>2</sub> 0Me	Ph C00-CH <sub>2</sub> -C00Bt
40	Compound	OH CONH	- CONH
45		MeN(CH <sub>2</sub> ) 40 C1 + C1	C1 H <sub>2</sub> N-(CH <sub>2</sub> ), -0 —C1
50	Ex. No.	MeN(CF H 42 • HC1	43 •HC1

5		Blemental analysis (%)	C <sub>2</sub> 7H <sub>3</sub> Cl <sub>2</sub> N <sub>2</sub> O <sub>7</sub> • HCl Calculated C, 53, 52 H, 5, 82 N, 4, 62 Found C, 53, 34 H, 5, 97 N, 4, 39	C2.644.2C1.2N208.4C1 Calculated C. 55.77 H. 5. 94 N. 5. 00 Found C. 55. 37 H. 6. 02 N. 4. 86
15		PAB-NS	569 (free base, MH+)	523 (free base, MH+)
20		IR (cm <sup>-1</sup> )	Neat 2971 1754 1640 1584 1460	KBr 3422 2939 1718 1641 1585 1458
<i>25</i>	Table 24	'H-NMR & (ppm), 300MHz	DNSO-de 1. 11 (9H, s) 1. 77-1. 91 (4H, m) 2. 54 (3H, s) 2. 75-3. 25 (4H, m) 4. 00-4. 10 (2H, m) 4. 40-4. 80 (1H, m) 5. 76 (2H, s) 7. 20-7. 40 (5H, m) 8. 17 (1H, s) 8. 74 (2H, brs) 9. 55 (1H, brs) 13. 29 (1H, s)	DMSO-d <sub>6</sub> 1. 15-1. 90(14H, m) 2. 82-2. 93(2H, m) 3. 10-3. 24(2H, m) 4. 01-4. 08(2H, m) 4. 65-4. 75(2H, m) 7. 18-7. 32(5H, m) 7. 92(3H, brs) 8. 21(1H, s) 9. 47(1H, d) 13. 39(1H, brs)
35			Ph C00CH20C0 ——	$\bigcirc$
40		Compound	OH COUNTY COU	OH CONH
45			C1 MeN-(CH <sub>2</sub> ), 4-0-(C1 -HC1	C1 H <sub>3</sub> N-(CH <sub>2</sub> ), -0 —(C1
50		Ex. No.	Meh 1 •	H <sub>2</sub> N

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			·	
5		Elemental analysis (%)	C <sub>27</sub> H <sub>34</sub> Cl <sub>2</sub> N <sub>2</sub> O <sub>6</sub> ·HCl Calculated C, 56.50 H, 6.15 N, 4.88 Found C, 54.51 H, 5.63 N, 4.64	
15		FAB-MS	537 (free base, MH+)	free base.
20	-	IR (cm <sup>-1</sup> )	KBr 2938 1641 1584 1458 1357 1219	KBr 1718 1718 1642 1584 1221
25	Table 25	'H-NNR & (ppm), 300MHz	DMSO-d <sub>6</sub> 1. 13-1. 92(10H, m) 2. 55(3H, t, J=6. 0Hz) 2. 85-3. 02(2H, m) 3. 08-3. 26(2H, m) 4. 00-4. 11(2H, m) 7. 25-7. 34(5H, m) 8. 20(1H, s) 8. 64(2H, brs) 9. 43(1H, d, J=6. 0Hz) 13. 39(1H, s)	DMSO-d <sub>8</sub> 0. 66-0. 88(4H m) 0. 92-1. 92(8H m) 2. 50(6H, d, J=3. 0Hz) 2. 72-2. 92(2H, m) 3. 22-3. 78(4H, m) 4. 04-4. 12(2H, m) 7. 22-7. 42(6H, m) 8. 20(1H, s) 9. 44((1H, br) 13. 43(1H, s)
30		H <sub>1</sub>	9.9.6.4.4.0.00.0.E.	0.004444.000.E
35			Ph C00	Ph Me Coo Me
40		Compound	OH CONH	OH
45			C1 MeN-(CH <sub>2</sub> ),4-0 H •HC1	C1)
50		Bx. No.	W 46	Н 47

5	Elemental analysis (%)	·	
10	FAB-MS	538 (free base, MH+)	575 (free base. MH*)
15	IR (cm <sup>-1</sup> )	Neat 2964 1740 1674 1584 1458	KBr 3386 2909 1718 1642 1585 1541
	m), 300MHz		(2H, m) (2H, m) (2H, m) (2H, m) (2H, m) (2H, m) (5H, m) (5H, m) (5H, m) (5H, m) (5H, m)
rable 26	1H-NMR & (ppm), 300MHz	DMSO-d <sub>6</sub> 1, 60-2, 15(8H, m) 2, 72(3H, s) 2, 80-3, 60(6H, m) 4, 05-4, 10(2H, m) 4, 60-4, 91(2H, m) 7, 20-7, 39(6H, m) 7, 82(3H, brs) 8, 19-8, 26(1H, m)	DMSO-de 1. 40-1. 53(2H 1. 65-1. 96(16) 2. 82-2. 93(2H 3. 13-3. 40(2H 4. 05(2H, t. J= 4. 74-4. 88(2H 7. 08-7. 16(5H 7. 91(3H, brs) 8. 21(1H, s) 9. 48(1H, d. J=
30		N-We	
35	pu	H-083	4 000
40	Compound	O CON	OH
45		C1 H <sub>2</sub> N-(CH <sub>2</sub> ), 4-0	CI CH2),4-0 —
50	<u> </u>	H <sub>2</sub> N .	H2N-()

8x.

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!			
5	Blemental analysis (%)		<u>.</u>
10	PAB-MS	512 (free base, MH*)	526 (free base, MH*)
15	IR (cm <sup>-1</sup> )	KBr 3398 2958 1736 1641 1585 1542	Neat 2951 1747 1661 1556
20			. ^
rable 27	'H-NMR & (ppm), 300MHz	DMSO-d <sub>4</sub> 1. 75-1. 91 (4H, m) 2. 54 (3H, t, J=4. 5Hz) 2. 90-3. 08 (3H, m) 3. 22 (1H, dd, J=12. 3Hz) 3. 82 (2H, d, J=6Hz) 3. 98-4. 08 (2H, m) 4. 78-4. 88 (1H, m) 7. 12-7. 37 (5H, m) 8. 27 (1H, s) 8. 64 (1H, t, J=6Hz) 9. 30 (1H, d, J=6Hz) 13. 20 (1H, brs) 13. 52 (1H, s)	DMSO-d <sub>4</sub> 1. 77-1. 91(4H, m) 2. 54(3H, t, J=6Hz) 2. 95(2H, brs) 3. 03(1H, dd, J=15, 12Hz) 3. 22(1H, dd, J=15, 6Hz) 3. 91(2H, d, J=6Hz) 4. 00-4. 09(2H, m) 4. 77-4. 87(1H, m) 7. 13-7. 38(5H, m) 8. 27(1H, s) 8. 77-8. 87(3H, m) 9. 34(1H, d, J=9Hz) 13. 52(1H, s)
30		H	<u>o</u>
. 35		CONHCH <sub>2</sub> COOH	CONHICH 2 COOMe
40	Compound	OH	HO
<b>45</b>		CI MeN-(CH <sub>2</sub> ),-0 — H •HCl	C1.  MeN-(CH <sub>2</sub> ),-0  H  C1.
50	Bx. No.	50	51

5	Elemental analysis (%)	Ca2Ha,Cl2N104S·HCl Calculated C, 50.63 H, 5.21 N, 5.37 Pound C, 50.40 H, 5.29 N, 5.28	CarHs*ClsN208·HCl Calculated C. 57. 11 H. 5. 15 N. 4. 93 Found C. 56. 97 H. 5. 22 N. 5. 15
15	FAB-MS	485 (free base. MH*)	531 (free base, MH+)
20	[R (cm <sup>-1</sup> ).	KBr 2930 1641 1584 1535 1457 1226	KBr 3397 2958 1719 1642 1586 1543
Table 28	'H-NMR & (ppm), 300Mfz	DMSO-ds 1. 17(3H, t, J=6. 0Hz) 1. 62-1. 92(4H, m) 2. 77-2. 97(4H, m) 3. 09(1H, dd, J=15. 0. 12. 0Hz) 3. 11-3. 35(1H, m) 4. 82-4. 96(1H, m) 7. 13-7. 36(5H, m) 7. 13-7. 36(5H, m) 7. 92(2H, brs) 8. 25(1H, s) 9. 61-9. 73(1H, m) 13. 23(1H, s)	DNSO-d <sub>4</sub> 1. 82(4H, m) 2. 80(2H, m) 3. 16(1H, dd, J=9, 12Hz) 3. 24(1H, dd, J=6, 12Hz) 4. 05(2H, brs) 4. 81(1H, ddd, J=6, 7, 9Hz) 5. 14(1H, d, J=12Hz) 5. 17(1H, d, J=12Hz) 7. 16-7. 39(10H, m) 7. 16-7. 39(10H, m) 7. 91(3H, brs) 8. 19(1H, s) 9. 50(1H, s) 13. 32(1H, s)
30	#	DNSO-d 1. 17(3H 1. 62-1. 2. 77-2. 3. 09(1H 3. 11-3. 3. 99-4. 4. 82-4. 7. 13-7. 7. 92(2H 8. 25(1H 9. 61-9.	
35	-	Ph Coset	- Ph C00CH2Ph
40	Compound	HOCOM	OH CONH -
45		C1 C1-	C1 H <sub>2</sub> N-(CH <sub>2</sub> ),-0 — -HC1
50	Ex. No.	H <sub>2</sub> N	F3 .

. 5	Blemental analysis (%)	CasHaoClaNaOs·HCl Calculated C, 57, 79 H, 5, 37 N, 4, 81 Pound C, 57, 34 H, 5, 44 N, 4, 78	C <sub>2.8</sub> H <sub>3.2</sub> Cl <sub>2</sub> N <sub>2</sub> O <sub>8.</sub> HCl Calculated C, 58.45 H, 5.58 N, 4.70 C, 58.18 H, 5.49 N, 4.72
15	PAB-MS	545 (free base, MH <sup>+</sup> )	558 (free base, MH+)
20	IR (cm <sup>-1</sup> )	KBr 3412 3300 2958 2789 1745 1639 1584 1541	KBr 2957 2690 1740 1638 1584 1456
rable 29	'H-NMR & (ppm), 300MHz	0-d <sub>4</sub> (4H, brs) (8H, t, J=6Hz) (2H, brs) (1H, d, J=10, 12Hz) (1H, d, J=6, 12Hz) (2H, brs) (2H, ddd, J=6, 7, 9Hz) (1H, d, J=12Hz) (1H, d, J=7Hz) (2H, brs) (2H, brs)	DMSO-d <sub>4</sub> 1. 76-1. 95(4H, m) 2. 75(6H, s) 3. 05-3. 30(4H, m) 4. 06(2H, t, J=7Hz) 4. 75-4. 87(1H, m) 5. 10-5. 20(2H, m) 7. 18-7. 40(10H, m) 8. 18(1H, s) 9. 52(1H, brs) 10. 20(1H, brs) 13. 40(1H, brs)
30	IN-H1	DMSO 2. 2. 44(0) 2. 3. 2. 16(0) 3. 2. 16(0) 3. 2. 16(0) 4. 4. 0. 17(0) 5. 17(0) 9. 50(0) 13. 32	DMS0 1. 76(-) 3. 05-(-) 5. 10-4 7. 18-7 7. 18-7 10. 20 13. 40
35		Ph COOCH2Ph	Ph C00CH <sub>2</sub> Ph
40	Compound	OH	OH CONH
	3	MeN-(CH <sub>2</sub> ), -0 C1 H C1	Me <sub>2</sub> N-(CH <sub>2</sub> ),-0—C1
	8. 8.	ى 2.	55 55

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5		Elemental analysis (%)	C2.6H3.C12N4.06.3HC1 Calculated C. 47.11 H. 5.63 N. 8.45 Found C. 45.84 H. 5.72 N. 7.76	
			9.0	.se,
15		FAB-MS	553 (free base, MH*)	536 (free base, MH+)
20		(cm <sup>-1</sup> )	KBr 3423 2957 1751 1638 1585 1542 1458	KBr 3422 2937 1752 1639 1584 1541 1457 1346
25	Table 30	'H-NMR & (ppm), 300MHz	DMSO-d <sub>6</sub> 1. 82(4H, m) 2. 95(2H, m) 2. 95(2H, m) 3. 08-3. 55(12H, m) 4. 04(2H, brs) 4. 41(2H, m) 7. 12-7. 36(5H, m) 7. 97(3H, brs) 8. 31(1H, s) 9. 63(3H, m)	DMSO-de 1. 42-1. 86(10H, m) 2. 22-2. 40(4H, m) 2. 72-2. 84(2H, m) 3. 18-3. 28(2H, m) 4. 66-4. 72(1H, m) 7. 16-7. 34(5H, m) 7. 84(2H, br) 8. 20. 8. 22(1H, s) 9. 34. 9. 53 (1H, d, J=5. 8H2) 13. 32. 13. 48
30			( <del>Z</del> )	$\wedge$
35		P	Ph C00(CH <sub>2</sub> ) <sub>2</sub> -N NH	Ph-000-N
40		Compound	OH	OH CONH
<b>4</b> 5			C1 C1 C1 C1	C1 H2N-(CH2), -0
50	-			
		S.S.	56	57.

			Table 31		
Compound	q		'H-NAIR & (ppm), 300MHz	IR (cm <sup>-1</sup> )	Œ.
15	5	/Ph	DMS0-de	KBr	വ

Bx. No.	Compound	'H-NAIR & (ppm), 300MHz	1R (cm <sup>-1</sup> )	FAB-MS	Elemental analysis (%)
28	Me-N N-(CH <sub>2</sub> ) <sub>2</sub> -0 CONH COOMe  •2HC1	DMSO-de 2. 81(3H, s) 3. 13-3. 80(10H, m) 4. 27-4. 47(2H, m) 4. 66-4. 83(1H, m) 7. 13-7. 32(5H, m) 8. 22(1H, s) 9. 49(1H, d, J=8. 5Hz) 13. 71(1H, brs)	KBr 1740 1641 1584 1457 1355 1220	510 (free base, MH+)	Ca, Hz, Cl, N, O, · 2HCl Calculated C, 49, 42 H, 5, 36 N, 7, 20 Pound C, 47, 94 H, 5, 52 N, 6, 77
29	HN N-(CH <sub>2</sub> ) <sub>2</sub> -0 -CONH -COOMe	DMSO-d <sub>6</sub> 1. 15(3H. d. J=6. 2Hz) 2. 78-3. 95(13H. m) 3. 67(3H. s) 4. 54-4. 72(1H. m) 7. 17-7. 34(5H. m) 7. 45(1H. s) 8. 69-8. 82(1H. m) 12. 27-12. 36(1H. m)		510(MH+)	

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5		Elemental analysis (%)		Optical rotation: [α] <sup>15</sup> <sub>b</sub> = -53.0° (c=0.37, MeOH)
15		FAB-MS	510 (free base, MH+)	510 (free base. MH+)
20		IR (cm <sup>-1</sup> )		KBr 3427 1736 1641 1458 1222
25	Table 32	'H-NMR & (ppm), 300MHz	DMSO-d <sub>4</sub> 1. 30(3H, d, J=6Hz) 3. 00-3. 70(11H, m) 3. 67(3H, s) 4. 42(2H, s) 7. 18-7. 30(5H, m) 8. 23(1H, s) 9. 52(1H, d, J=9Hz) 9. 80(1H, br) 13. 39(1H, br)	DMSO-d <sub>6</sub> 1. 29(3H, d, J=6. 3Hz) 3. 00-3. 20(9H, m) 3. 66(3H, s) 4. 41(2H, brs) 4. 77(1H, m) 7. 15-7. 30(5H, m) 8. 22(1H, s) 9. 49(1H, d, J=7. 6Hz) 9. 70(2H, br) 13. 35(1H, brs)
35			- Ph COOMe	COOME
40		Compound	C1 OH -CONH -	C1 OH CONH -
			HN Ne -2HC1	Hy N-(CH <sub>2</sub> ) Nè
50		Bx. No.	09	61

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5	Elemental analysis (%)	C <sub>24</sub> H <sub>2</sub> ,Cl <sub>2</sub> N <sub>3</sub> O <sub>6</sub> ·2HCl Calculated C, 54, 16 H, 5, 13 N, 6, 11 Pound C, 53, 21 H, 5, 25 N, 5, 96	
15	PAB-MS	510 (free base, MH+)	524 (free base, MH*)
20	IR (cm <sup>-1</sup> )	KBr 3425 2450 1747 1664 1452 1248 1213	
Table 33	'H-NMR & (ppm), 300MHz	DMSO-d <sub>6</sub> 1. 30(3H, d, J=6. 0Hz) 3. 09-3. 83(11H, m) 3. 67(3H, s) 4. 34-4. 47(2H, m) 4. 73-6. 81(1H, m) 7. 17-7. 29(5H, m) 8. 23(1H, s) 9. 51(1H, d, J=6. 0Hz) 9. 63-9. 92(1H, m) 13. 35-13. 47(1H, m)	DMSO-d <sub>4</sub> 1. 34(3H, d, J=8Hz) 2. 80(3H, s) 3. 00-3. 70(11H, m) 3. 67(3H, s) 4. 37(2H, brs) 4. 75(1H, m) 7. 15-7. 32(5H, m) 8. 22(1H, s) 9. 50(1H, d, J=6Hz) 13. 39(1H, s)
35		COOMe	H COOMe
40	Compound	C1 OH CONH -	10 -0- 0- 10
45	-	-CH <sub>2</sub> ) <sub>2</sub> -C	Me N-(CH2)2
50	Ex. No.	Me 62	Me 63

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	8		
5	Elemental analysis		
15	FAB-MS	510 (free base, MH*)	524 (MH+)
20	IR (cm <sup>-1</sup> )	i .	KBr 3422 2940 2360 1736 1638 1456
	Table 34 "H-NMR & (ppm), 300MHz	DMS0-d <sub>6</sub> 1. 14(3H, t, J=7. 6Hz) 2. 48-2. 52(8H, m) 3. 14-3. 38(4H, m) 4. 11(2H, q, J=7. 6Hz) 4. 39(2H, bs) 4. 68-4. 80(1H, m) 7. 18-7. 32(5H, m) 8. 23(1H, s) 9. 47(1H, d, J=8. 8Hz)	ASO-d <sub>6</sub> 11(3H, t, J=6.0Hz) 50-3.20(13H, m) 10-4.10(4H, m) 52-4.66(1H, m) 20-7.30(5H, m) 75(1H, s)
30	=	51.48.444.88	Et 2. (
35		NH COOBt	CONH COO
40	Compound	10 0- 0- 10	10 C1
45		C1 HN N-(CH <sub>2</sub> ) <sub>2</sub> -0 -	C1 Me-N N-(CH <sub>2</sub> ) <sub>2</sub> -0 C1
			<del></del>

5	Elemental analysis	C1.6H2.1C1.2N3.04.2HC1 Calculated C, 50. 27 H, 5. 57 N, 7. 03 Pound C, 49. 88 H, 5. 56 N, 6. 93	C2.6H3.C12N3O6.2HC1 Calculated C, 50.27 H, 5.57 N, 7.03 Pound C, 49.68 H, 5.68 N, 6.66
. 15	FAB-MS	524 (free base, MH*)	524 (free base, MH+)
20	IR (cm <sup>-1</sup> )	KBr 3423 1740 1640 1584 1356 1219	KBr 2361 2343 1584 1458 1352 1216
25 5 	Table 35 H-NMR & (ppm), 300MHz	DMSO-d <sub>6</sub> 2. 10-2. 35(2H, m) 2. 80(3H, s) 3. 10-3. 94(12H, m) 3. 66(3H, s) 4. 10-4. 22(2H, m) 4. 62-4. 72(1H, m) 7. 20-7. 41(5H, m) 8. 20(1H, s) 9. 47(1H, d, J=6. 0Hz) 13. 3(1H, br s)	DMSO-d <sub>4</sub> 1. 14(3H, t, J=6. 0Hz) 2. 23(2H, m) 3. 00-3. 85(10H, m) 4. 05-4. 16(4H, m) 4. 67-4. 77(1H, m) 7. 12-7. 35(5H, m) 7. 50-7. 66(1H, m) 8. 21(1H, brs) 9. 40-9. 60(1H, brs) 9. 45(1H, d, J=6. 0Hz) 13. 40(1H, brs)
<b>35</b>	-	Ph 22 33 33 34 4 4 4 4 4 4 4 4 4 4 4 4 4 4	Ph C00Et 3
40 .	Compound	CI OH CONE	CI CONH
45		Me-N N-(CH <sub>2</sub> ).	.2HC1
50	RX.		- F9

5 10	Elemental analysis (%)	·	
15	FAB-MS	538 (free base, MH <sup>+</sup> )	509 (free base, M <sup>†</sup> )
	IR (cm <sup>-1</sup> )		KBr 3372 2940 2805 2726 2489 1739 1642 1585 11460 11412 1352
20	300MHz		2.0Hz)
	Table 36 <sup>1</sup> H-NMR & (ppm), 300MHz	DMSO-46 1.21(3H, t, J=8Hz) 2.25(2H, brs) 2.82(3H, s) 3.08-3.90(12H, m) 4.06-4.13(4H, m) 4.70(1H, m) 7.16-7.28(5H, m) 8.21(1H, s) 9.45(1H, d, J=8Hz) 13.36(1H, brs)	DMSO-d <sub>6</sub> 1.12(3H, t, J=6.9Hz) 1.29-1.47(2H, m) 1.73(2H, dd, J=5.7, 12.0Hz) 1.80-1.95(4H, m) 2.85(2H, m) 3.07-3.28(4H, m) 4.04-4.13(4H, m) 7.16-7.28(5H, m) 7.16-7.28(5H, m) 8.20(1H, s) 8.65(1H, brs) 9.50(1H, brs) 13.35(1H, brs)
	I.	-Ph DMSO-d <sub>6</sub> 1.21(3H, t, J= -C008t 2.25(2H, brs) 2.82(3H, s) 3.08-3.90(12I) 4.06-4.13(4H, m) 7.16-7.28(5H, s) 8.21(1H, s) 9.45(1H, d, J- 13.36(1H, brs)	-Ph DMSO-46 1.12(3H, t 1.12(3H, t 1.73(2H, d 1.80-1.95(2H, n 3.07-3.28( 4.04-4.13( 4.67-4.75( 7.16-7.28(1H, b 8.20(1H, b 8.87(1H, b 9.50(1H, b
35		HNOO	CONH
40	Compound	15	10
45		Me-N N-(CH2).	HC1
50	BX.	Me-	H- 69

	Elemental analysis (%)		C <sub>23</sub> H <sub>30</sub> Cl <sub>2</sub> N <sub>2</sub> O <sub>5</sub> ·HCl Calculated C, 56.91 H, 6.23 N, 5.77 Found C, 56.90 H, 6.29 N, 5.73
15	PAB-MS	523 (free base, MH <sup>+</sup> )	449.1 (free base, MH <sup>+</sup> )
	IR (cm <sup>-1</sup> )	KBr 3406 2938 1736 1638 1584 1460 1412 1352 1215 1075 701	
20	2	<b>(</b> 2)	
7aple 37	¹H-NMR & (ppm), 300MHz	DMSO-4 <sub>6</sub> 1.23(3H, t, J=7.1Hz) 1.42-1.57(2H, m) 1.68-1.98(5H, m) 2.69(3H, s) 2.88-2.96(2H, m) 3.10-3.24(2H, m) 3.31-3.39(2H, m) 4.05-4.14(4H, m) 4.72(1H, ddd, J=6.3, 7.5, 9.1Hz) 7.15-7.29(5H, m) 8.21(1H, s) 9.48(1H, brs) 13.35(1H, brs)	CDCl <sub>3</sub> 1.13(3H, t, J=7.0Hz) 1.70-1.88(4H, m) 2.49-2.53(5H, m) 3.07-3.21(2H, m) 4.06-4.13(4H, m) 4.69(1H, dd, J=8.4, 15.6Hz) 6.67(1H, s) 7.18-7.32(5H, m) 8.04(1H, s) 8.75(1H, brs) 8.99(1H, d, J=7.2Hz) 12.51(1H, brs)
30	1H-1	SO-d (33 + 1.5) (31 + 1.6) (31 +	Cl3 3(3H) 2-2.5 7-3.2 7-3.2 7(1H) 7(1H) 7(1H) 7(1H)
35 40 45	Compound	HC1 CI OH Ph DMSO-d <sub>6</sub> 1.23(3H, t <sub>1</sub> 1.42-1.57(2 1.68-1.98(5 2.69(3H, s) 2.69(3H, s) 2.69(3H, s) 2.69(3H, s) 2.69(3H, s) 2.69(3H, d) 2.69(3H, d) 2.16-7.29(5 8.21(1H, s) 9.48(1H, d) 10.34(1H, b) 13.35(1H, b)	MeN-(CH <sub>2</sub> ), -0 CONH CDC! <sub>3</sub> 1.13(3H, t, 1
50	36	70	
	S.S.	1.	

5	Elemental analysis (%)		
15	FAB-MS	463 (free base, MH <sup>+</sup> )	479 (free base, MH <sup>+</sup> )
	IR (cm <sup>-1</sup> )	KBr 3428 2958 2686 1736 1604 1541 1493 1375 1267	KBr 1741 1637 1489 1265
20	, 300MHz	4.4Hz)	
Table 38	<sup>1</sup> H-NMR & (ppm), 300MHz	DMSO-46 1.13(3H, t, J=7.0Hz) 1.75-1.90(4H,brs) 2.75(3H, s) 2.75(3H, s) 3.12-3.33(4H, m) 4.10(4H, m) 4.0(1H, dd, J=5.4, 14.4Hz) 6.65(1H, brs) 7.21-7.29(5H, m) 8.04(1H, s) 8.98(1H, d, J=7.8Hz) 12.51(1H, s)	CDCl <sub>3</sub> 1.95-2.10(4H, m) 2.70(3H, s) 3.11(2H, t, J=7.5Hz) 3.21(2H, m) 3.77(3H, s) 4.01(2H, t, J=6.0Hz) 5.00(1H, m) 6.38(1H, s) 7.02(1H, d, J=7.2Hz) 7.14-7.32(5H, m) 7.52(1H, s) 9.43(2H, brs) 12.2(1H, s)
30	H <sub>1</sub>		CDCl <sub>3</sub> 1.95-2.10(4H 2.70(3H, s) 3.11(2H, t, Ji 3.21(2H, m) 3.77(3H, s) 4.01(2H, t, Ji 5.00(1H, m) 6.38(1H, s) 7.02(1H, d, Ji 7.14-7.32(5H 7.52(1H, s) 9.43(2H, brs) 12.2(1H, s)
35		Ph COORE	COOMe
40	Compound	OH CONH	OH CONH -
45		Me <sub>2</sub> N-(CH <sub>3</sub> ),-0 — Cl <sup>2</sup>	MeN-(CH.),-0— H •HCl
50	Bx. No.	Me <sub>2</sub> N-	MeN-( H -HC1

		<del></del>	
5	Elemental analysis (%)	C <sub>22</sub> H <sub>27</sub> BrN <sub>2</sub> O <sub>5</sub> ·HCl Calculated C, 51.23 H, 5.47 N, 5.43 Pound C, 50.93 H, 5.51 N, 5.34	
15	FAB-MS	479 (free base, MH <sup>+</sup> )	493 (free base, MH <sup>+</sup> )
	IR (cm <sup>-1</sup> )	KBr 1736 1601 1489 1373 1263	KBr 3374 2960 1736 1638 1599 1376 1199
20	300MHz	·	_
rable 39	'H-NMR & (ppm), 300MHz	DMSO-d <sub>6</sub> 1.13(3H, t, J=7.0Hz) 1.70-1.89(4H, m) 2.87(2H, m) 3.2(2H, m) 4.0-4.1(4H, m) 4.7(1H, m) 6.65(1H, s) 7.15-7.30(5H, m) 7.92(3H, brs) 8.17(1H, s) 8.98(1H, d, J=5.6Hz) 12.51(1H, s)	CDCl <sub>3</sub> 1.27(3H, t, 1=7.2Hz) 1.93-2.02(2H, m) 2.07-2.17(2H, m) 2.71(3H, s) 3.1-3.2(2H, m) 3.2-3.3(2H, m) 4.03(2H, t, 1=6Hz) 4.03(2H, t, 1=6Hz) 4.23(2H, g, 1=7Hz) 4.98(1H, dt, 1=7, 6Hz) 6.88(1H, d, 1=7, 8Hz) 7.14-7.18(2H, m) 7.22-7.32(3H, m) 7.22-7.32(3H, m) 7.22-7.32(3H, s) 9.51(2H, brs) 12.29(1H, brs)
30	1 <sup>1</sup> H-	DMSO-d <sub>6</sub> 1.13(3H, t, J= 1.70-1.89(4H 2.87(2H, m) 3.2(2H, m) 4.0-4.1(4H, m) 6.65(1H, s) 7.15-7.30(5H 7.92(3H, brs) 8.98(1H, d, J) 12.51(1H, s)	CDCl <sub>3</sub> 1.27(3H, t, J= 1.93-2.02(2H, 2.07-2.17(2H, S) 3.1-3.2(2H, m) 3.2-3.3(2H, m) 4.03(2H, t, J= 4.23(2H, q, J= 4.23(2H, q, J= 6.39(1H, s) 6.88(1H, d, J= 7.14-7.18(2H, 7.22-7.32(3H, 7.49(1H, s) 9.51(2H, brs) 12.29(1H, brs)
35		Ph C00Bt	C00Bt
40	Compound	OH CONH	OH CONH
45		H±N-(CH2),-0 —- Br'	MeN-(CH1),-0-/-
50	Bx. No.	H 74	7.5 M

5	Elemental analysis (%)		
15	FAB-MS	449 (free base, M-1)	(free base, MH <sup>+</sup> )
	IR (cm <sup>-1</sup> )	·	
20	300MHz	·	
Table 40	H-NMR & (ppm), 300MHz	s s s s s s s s s s s s s s s s s s s	CDC3 1.13(3H, t, J=7Hz) 1.85-1.95(4H, m) 2.04(3H, s) 2.18(3H, s) 2.18(3H, m) 2.93(2H, m) 3.16-3.20(2H, m) 3.16-3.20(2H, m) 3.76(2H, m) 4.10(2H, q, J=7Hz) 4.69(1H, m) 7.17-7.32(5H, m) 7.71(1H, s) 9.01(2H, brs) 9.07(1H, d, J=8Hz) 12.66(1H, s)
30 E	N-H <sub>1</sub>	DMSO-d <sub>6</sub> 1.91(4H, m) 2.12(3H, s) 2.57(3H, s) 2.98(2H, m) 3.10-3.30(2H, m) 3.70(3H, s) 3.99(2H, m) 4.78(1H, m) 6.35(1H, s) 7.15-7.30(5H, m) 7.65(1H, s) 8.62(1H, brs) 9.24(2H, brs) 1.36(1H, brs)	CDCl <sub>3</sub> 1.13(3H, t, J=7H, 1.85-1.95(4H, m) 2.04(3H, s) 2.18(3H, s) 2.18(3H, m) 2.93(2H, m) 3.16-3.20(2H, m) 3.16-3.20(2H, m) 3.16(2H, m) 4.69(1H, m) 7.17-7.32(5H, m) 7.71(1H, s) 9.01(2H, brs) 9.07(1H, d, J=8H) 12.66(1H, s)
35		Ph C00Me	C00Et
40	Compound	OH	OH
45		MeN-(CH1),-0 — H Me'	MeN-(CH <sub>2</sub> ),-0— H H Me
50	BX. No.	M6 .	Ne
	1	i ·	

	_		
5		Blemental analysis (%)	<i>:</i>
15		IR (cm <sup>-1</sup> ) FAB-MS	415 (free base, MH <sup>+</sup> )
20		IR (cm <sup>-1</sup> )	
		ı), 300MHz	(2)
25 30	Table, 41	<sup>1</sup> H-NMR & (ppm), 300MHz	DMSO-d <sub>6</sub> 1.63-1.78(4H, m) 2.02(3H, s) 2.55(3H, s) 2.82-3.11(4H, m) 3.62(3H, s) 3.84-3.95(2H, m) 4.58-4.65(1H, m) 6.21(1H, d, J=2.4Hz) 6.24(1H, d, J=2.4Hz)
35			COOMe
40		Compound	OH CONH
45			N-(CH <sub>8</sub> ),-0-/HC1

			18.010			
mz	Š.	Compound	<sup>1</sup> H-NMR & (ppm), 300MHz	IR (cm <sup>-1</sup> )	FAB-MS	Blemental analysi (%)
		hQ HO	9p-OSWQ		415	
		(	1.63-1.78(4H, m)		(min com)	
<u>-</u>		Men-(CH <sub>2</sub> ) - 0 - 0 - 0 - 0 - 0 - 0 - 0 - 0 - 0 -	2.02(3H, s)		( HIM	
			2.55(3H; s)			
		Sec.	2.82-3.11(4H, m)			
_		104.	3.62(3H, s)			
	9		3.84-3.95(2H, m)			•
	0		4.58-4.65(1H, m)			
			6.21(1H, d, J=2.4Hz)			
			6.24(1H, d, J=2.4Hz)		-	
			7.18-7.36(5H, m)			-
			8.38(1H, d, J=7.5Hz)			
			8.62-8.78(2H, m)			
			9.70(1H, brs)			-
		40 10	DMSO-4 <sub>K</sub>		431	
			[1.80(4H. m)		(free base,	
_		MeN-(CH <sub>2</sub> ), -0 -(C) - CONH - COOMe	2.50(3H, m)		MH <sup>+</sup> )	
		入 ·	2.95(2H, m)			٠.
		,0°H0H	3.08-3.22(2H, m)			
			3.65(3H, s)			
	ć		3.95-4.15(2H, m)			
_	3		4.60-4.75(3H, m)			-
			6.54(1H, s)			
			7.15-7.32(5H, m)			
			7.99(1H, s)			
<u>.</u>			8.68(2H, brs)			
			8.93(1H, d, J=8Hz)			
			12.54(1H, s)			

5	Elemental analysis (%)		
15	FAB-MS	493 (free base, MH <sup>+</sup> )	(free base, MH <sup>†</sup> )
	IR (cm <sup>-1</sup> )		
20	, 300MHz		
7a 1a 42	<sup>1</sup> H-NMR & (ppm), 300MHz	DMSO-46 1.71-1.83(4H, m) 2.02(3H, s) 2.55(3H, s) 2.88-3.11(4H, m) 3.63(3H, s) 3.92-4.03(1H, m) 6.45(1H, s) 8.57-8.78(2H, m) 8.57-8.78(2H, m) 8.57-8.78(2H, m)	DMSO-d <sub>6</sub> 1.79-1.93(4H, m) 2.00(3H, s) 2.00(3H, s) 2.91-3.11(4H, m) 3.63(3H, s) 3.83-3.95(2H, m) 4.60-4.66(1H, m) 7.20-7.36(5H, m) 8.54-8.66(2H, m) 8.54-8.66(2H, m) 8.54-8.66(1H, d, J=7.8Hz)
30	11.	DMSO-d <sub>6</sub> 1.71-1.83(4H, 2.02(3H, s) 2.55(3H, s) 2.55(3H, s) 2.88-3.11(4H, 3.63(3H, s) 3.92-4.03(1H, s) 6.45(1H, s) 7.08-7.37(5H, g) 8.57-8.78(2H, g) 9.60(14, d, J, g)	DMSO-d <sub>6</sub> 1.79-1.93(4) 2.00(3H, s) 2.56(3H, s) 2.56(3H, s) 2.91-3.11(4) 3.63(3H, s) 3.83-3.95(2) 4.60-4.66(1) 7.20-7.36(5) 8.54-8.66(2) 8.54-8.66(2) 8.93(1H, d, g) 9.61(1H, s)
35		COOMe	COOMe
40	Compound	Ne Ne	OH CONH
45		MeN-(CH <sub>1</sub> ),-0—( H Br	Br MeN-(CH <sub>2</sub> ),-0— H Br
50	Bx. No.	80 .	81 8

	(0)	T	-,
5	Blemental analysis (%)	C <sub>25</sub> H <sub>30</sub> Cl <sub>2</sub> N <sub>2</sub> O <sub>7</sub> ·HCl Calculated C, 51.96 H, 5.41 N, 4.85 Pound C, 51.88 H, 5.40 N, 4.82	C <sub>30</sub> H <sub>33</sub> O <sub>7</sub> N <sub>2</sub> O <sub>3</sub> ·HCI Ca1culated C, 56.31 H, 5.20 N, 4.38 Found C, 54.81 H, 5.30 N, 4.34
	<u>رم</u> ا	υ˙-	
15	FAB-MS	541 (free base, MH <sup>+</sup> )	603 (free base, M <sup>+</sup> )
20	IR (cm <sup>-1</sup> )	KBr 2961 1750 1461 1178	KBr 3426 2960 1717 1641 1664 1457 1278 1162
20	, 300MHz		.1Hz)
rable 43	<sup>1</sup> H-NMR & (ppm), 300MHz	DMSO-d <sub>6</sub> 1.19(3H, t, J=7.1Hz) 1.83(4H, brs) 2.94(2H, brs) 3.16-3.34(5H, m) 4.02-4.18(4H, m) 4.76(2H, d, J=2.2Hz) 4.84-4.92(1H, m) 7.16-7.36(5H, m) 8.22(1H, s) 8.87(2H, brs) 9.56(1H, d, J=7.2Hz) 9.56(1H, d, J=7.2Hz)	DMSO-d <sub>6</sub> 1.32(3H, t, J=7.0Hz) 1.82(4H, brs) 2.57(3H, brs) 2.97(2H, brs) 4.06(2H, brs) 4.32(2H, q, J=7.0Hz) 4.32(2H, dd, J=5.7, 14.1Hz) 7.16-7.36(7H, m) 8.01(2H, d, J=8.7Hz) 8.23(1H, brs) 8.33(2H, brs) 9.66(1H, brs) 11.3(1H, brs)
30	V-H <sub>I</sub>		
35		С00СН, С00Вц	-th -cog ————————————————————————————————————
40	Compound	OH CONH	OH CONH
45		C1 CH2), -0 C1 CH2)	-10 H CH <sub>2</sub> ),-0
50			3
	Bx.	83	83

5	Elemental analysis (%)		
15	FAB-MS	568 (free base, MH <sup>+</sup> )	582 (free base, MH <sup>+</sup> )
	IR (cm <sup>-1</sup> )		
20	, 300MHz		
rable 44	<sup>1</sup> H-NMR & (ppm), 300MHz	DMSO-d <sub>6</sub> 1.15(6H, t, J=7Hz) 1.75-2.00(6H, m) 2.49(3H, s) 2.90-3.05(8H, m) 3.10-3.25(2H, m) 4.00-4.21(4H, m) 7.17-7.30(5H, m) 8.28(1H, s) 8.88(2H, brs) 9.64(1H, brs) 13.37(1H, brs)	DMSO-d <sub>6</sub> 1.28(12H, m) 1.84(4H, m) 2.53(3H, t, J=5.7Hz) 2.94(2H, m) 3.13-3.93(4H, m) 3.63(2H, m) 4.01(2H, m) 4.37-4.48(1H, m) 7.18-7.29(5H, m) 8.31(1H, s) 8.86(2H, brs) 9.70(1H, brs) 10.03(1H, brs)
30	H <sub>1</sub>	DMSO-d <sub>6</sub> 1.15(6H, t, J=7 1.75-2.00(6H, 2.49(3H, s) 2.49(3H, s) 2.90-3.05(8H, 3.10-3.25(2H, 4.00-4.21(4H, m) 7.17-7.30(5H, s) 8.28(1H, s) 9.64(1H, brs) 10.43(1H, brs) 13.377(11, brs)	DMSO-d <sub>6</sub> 1.28(12H, m) 1.84(4H, m) 2.53(3H, t, J= 2.94(2H, m) 3.13-3.93(4H, m) 4.01(2H, m) 4.37-4.48(1H, r) 8.31(1H, s) 8.86(2H, brs) 10.03(1H, brs) 13.37(1H, brs)
35		-Ph -C00(CH <sub>2</sub> ),NBt <sub>2</sub>	-C00(CH <sub>2</sub> )
40	Compound	HOO3-	HO HO
45		MeN-(CH <sub>2</sub> ),-0-(CH <sub>2</sub> )	MeN-(CH <sub>2</sub> ),-0-(CH <sub>2</sub> )
50	Ex. No.	-2HCl	MeN-(C) H -2HC1
	L		<u></u>

5	Blemental analysis (%)		
15	FAB-MS	610 (free base, MH <sup>+</sup> )	455 (free base, MH <sup>+</sup> )
20	IR (cm <sup>-1</sup> )		
45 45	<sup>1</sup> H-NMR & (ppm), 300MHz	(E)	m) m1, =14, 6Hz) =14, 6Hz) Hz) Hz) =8, 6, 6Hz) m) m)
s Table 45	<sup>1</sup> H-NMR 8	Ph COO(CH <sub>2</sub> ) <sub>2</sub> N(CH <sub>2</sub> ) <sub>3</sub> Me 0.72-0.97(6H, m) (CH <sub>2</sub> ) <sub>3</sub> Me 1.51-1.73(4H, m) 1.75-1.83(4H, m) 2.83-3.15(6H, m) 3.16-3.39(5H, m) 4.02-4.10(2H, m) 4.75-4.91(1H, m) 7.20-7.30(5H, m) 8.36(1H, s) 8.68-8.96(2H, m) 10.56-10.73(1H, m) 10.56-10.73(1H, m) 13.32-13.47(1H, m)	CDCl <sub>3</sub> 1.80-1.89(2H, m) 1.91-2.01(2H, m) 3.21(1H, dd, J=14, 6Hz) 3.28(1H, dd, J=14, 6Hz) 3.76(2H, t, J=7Hz) 3.80(3H, s) 4.12(2H, t, J=6Hz) 5.03(1H, ddd, J=8, 6, 6Hz) 6.77(1H, d, J=8Hz) 7.09-7.13(2H, m) 7.28-7.35(3H, m) 12.64(1H, s)
35		12.) 2N(CH2.) 3Me (CH2.) 3Me	- Ph
40	Compound	<u> </u>	CONH HOO
<b>45</b>	Соп	C1 OH COM	HO-(CH <sub>2</sub> ), -0
50	Ex. No.	MAN-(CH <sub>2</sub> ), -0-(CH <sub>2</sub> ), -0-(C	но-(сн
	ωZ	∞	∞

5	Elemental analysis (%)		
15	FAB-MS	384 (free base, MH <sup>+</sup> )	399 (free base, MH <sup>†</sup> )
	IR (cm <sup>-1</sup> )		
20 .	300MHz		(Z)
rapie 46	<sup>1</sup> H-NMR & (ppm), 300MHz	DMSO-d <sub>6</sub> 1.70(2H, m) 1.85(2H, m) 2.04(2H, m) 2.62(3H, s) 2.93(2H, brs) 3.18-3.30(2H, m) 3.76(3H, s) 5.01(1H, m) 6.65(1H, d, J=7Hz) 6.78(1H, s) 9.35(2H, brs) 1.87(1H, s)	DMSO-d <sub>6</sub> 1.30(2H, m) 1.58(4H, m) 2.49(3H, s) 2.82(2H, t, J=8Hz) 3.11(1H, dd, J=14, 9Hz) 3.19(1H, dd, J=14, 6Hz) 3.65(3H, s) 4.74(1H, m) 6.75(2H, m) 7.18-7.31(5H, m) 7.81(1H, d, J=9Hz) 8.75(2H, brs) 9.00(1H, d, J=8Hz)
30 30	1.H.I	DMSO-d <sub>6</sub> 1.70(2H, m) 1.85(2H, m) 2.04(2H, m) 2.04(2H, s) 2.03(2H, s) 2.93(2H, s) 3.18-3.30(2H, s) 3.76(3H, s) 5.01(1H, m) 6.65(1H, s) 7.09-7.32(5H, s) 11.87(1H, s)	DMSO-de 1.30(2H, m) 1.58(4H, m) 2.49(3H, s) 2.82(2H, t, J- 3.11(1H, dd, 3.19(1H, dd, 3.65(3H, s) 4.74(1H, m) 6.75(2H, m) 7.18-7.31(5H) 7.81(1H, d, J- 8.75(2H, brs) 9.00(1H, d, J- 8.75(2H, brs)
35 ·		COOMe	Ph COOMe
40	Compound	OH CONH	OH CONH
45		MeN-(CH <sub>2</sub> ),—	MeN-(CH.), —-(
50	ex. No.	>≅ %	6

. [	sis				•								_	
	Blemental analysis (%)		•											•
	IR (cm <sup>-1</sup> ) FAB-MS	413	(rree base,	MH.										•
	IR (cm <sup>-1</sup> )													
Table 47	<sup>1</sup> H-NMR & (ppm), 300MHz	cDCl <sub>3</sub>	1.25-1,45(4H, m)	1.55-1.64(2H, m)	1.77-1.87(2H, m)	(2.54(2H, t, J=7.5Hz)	2.64(3H, s)	2.90(2H, t, J=7.8Hz)	3.23(2H, m)	6.63(2H, dd, J=8.1, 1.8Hz)	6.77(1H, d, J=1.5Hz)	6.88(2H, d, J=7.8Hz)	7,10-7,32(6H, m)	9.38(2H, brs)
	Compound	#9,		Men-(CH.)(C)-CONH // COOMe	)		Tou.							•
	RX.							S	2		•			

5	Elemental analysis (%)	C <sub>23</sub> H <sub>28</sub> Cl <sub>2</sub> N <sub>2</sub> O <sub>4</sub> ·HCl Calculated C, 54.83 H, 5.80 N, 5.56 Pound C, 54.63 H, 6.07 N, 5.48	
. 15	FAB-MS	467.0 (free base, MH <sup>+</sup> )	433 (frec base, MH <sup>†</sup> )
	IR (cm <sup>-1</sup> )	KBr 3429 2949 1743 1641 1587	
20	, 300MHz	5H2)	
79 6 148 75	H-NMR & (ppm), 300MHz	CDCl <sub>3</sub> 1.45-1.65(4H, m) 1.90-2.00(2H, m) 2.67(3H, s) 2.86-2.90(2H, t, J=7.5Hz) 2.80-3.05(2H, m) 3.18-3.31(2H, m) 3.80(3H, s) 7.15-7.32(6H, m) 9.48(2H, brs) 9.44(2H, s)	DMSO-46 1.36(2H, m) 1.45-1.65(4H, m) 2.49(3H, s) 2.62(2H, t, J=7Hz) 2.62(2H, m) 3.15(2H, m) 3.15(2H, m) 3.64(3H, s) 4.74(1H, m) 6.92(1H, s) 7.16-7.31(5H, m) 7.93(1H, s) 8.76(2H, brs) 9.05(1H, d, J=8Hz) 12.01(1H, s)
	1 <sub>B</sub>	CDCl <sub>3</sub> 1.45-1.65(4 1.90-2.00(2 2.86-2.90(2 2.80-3.05(2 3.18-3.31(2 3.18-3.31(2 3.18-3.31(2 7.15-7.32(6 7.15-7.32(6 1.248(2H, b)	DMSO-d <sub>6</sub> 1.36(2H, m) 1.45-1.65(4) 2.49(3H, s) 2.62(2H, t) 2.83(2H, m) 3.15(2H, m) 3.64(3H, s) 4.74(1H, m) 6.92(1H, s) 8.76(2H, ts) 8.76(2H, ts) 9.05(1H, s)
35		Ph COOMe	COOMe
40	Compound	OH	OH
45		MeN-(CH <sub>2</sub> ), Cl	Н НС1
50	Ex.	H H HeN-(	Meh 1.

5	Elemental analysis (%)		
. 15	FAB-MS	447 (free base, MH <sup>+</sup> )	461 (free base, MH <sup>+</sup> )
· .	IR (cm <sup>-1</sup> )	KBr 3422 2940 1738 1644 1644 1644 1107 1107 11096 11027	KBr 3423 2941 2693 1739 1644 1539 1483 1405 11212 1029 957 862 749
87 89 89 89	<sup>1</sup> H-NMR & (ppm), 300MHz	7.2Hz) m) m) m) m) m) m) m) m) 7.2Hz) 7.2Hz) J=6.2, 6.2, 7.5Hz) m)	CDC3 1.27(3H, t, J=7.1Hz) 1.34-1.46(2H, m) 1.61-1.71(2H, m) 1.83-1.94(2H, m) 2.69(2H, t, J=7.5Hz) 2.78(3H, s) 2.79(3H, s) 2.92-3.00(2H, m) 3.18-3.00(2H, m) 4.92(1H, dd, J=6.0, 6.0, 7.2Hz) 6.88(1H, s) 7.12-7.18(3H, m) 7.23-7.34(3H, m) 7.23-7.34(3H, m) 7.23-7.34(3H, m) 7.21-7.18(1H, brs) 11.60(1H, brs)
30	1H-NN	CDCl <sub>3</sub> 1.28(3H, t, J=7.2Hz) 1.40-1.51(2H, m) 1.57-1.67(2H, m) 1.84-1.95(2H, m) 2.62-2.68(5H, m) 2.88-3.01(2H, m) 3.16-3.29(2H, m) 4.24(2H, q, J=7.2Hz) 4.99(1H, ddd, J=6.2, 6.77(1H, m) 7.35(1H, s) 9.41(2H, brs) 11.67(1H, brs)	CDC3 1.27(3H, t, J=7.1Hz) 1.34-1.46(2H, m) 1.61-1.71(2H, m) 1.83-1.94(2H, m) 2.69(2H, t, J=7.5Hz) 2.78(3H, s) 2.78(3H, s) 2.92-3.00(2H, m) 3.18-3.30(2H, m) 4.22(2H, q, J=7.1Hz) 4.99(1H, ddd, J=6.0, 6.88(1H, s) 7.12-7.18(3H, m) 7.23-7.34(3H, m) 7.42(3H, s) 11.60(1H, brs) 12.26(1H, brs)
35		C00Bt	C00Bt
40	Compound	HO CONH	Ol
45		Men-(GH <sub>2</sub> ) s C1	Me <sub>2</sub> N-(CH <sub>2</sub> ) <sub>s</sub> C <sub>1</sub>
50	Ex. No.	-HC1	Me₂N •HC1

5	Blemental analysis (%)		
. 15	FAB-MS	477 (free base, MH <sup>+</sup> )	SS7 (free base, MH <sup>+</sup> )
. 20	IR (cm <sup>-1</sup> )	KBr 3375 1744 1641 1604 1540	
75 05 05 05	<sup>1</sup> H-NMR δ (ppm), 300MHz	CDCl <sub>3</sub> 1.4-1.6(4H, m) 1.90(2H, m) 2.50-2.66(5H, m) 2.94(2H, t, J=7.5Hz) 3.20(2H, m) 5.02(1H, s) 6.77(1H, s) 7.16-7.36(6H, m) 7.54(1H, bis) 11.57(1H, bis)	CDCl <sub>3</sub> 1.47-1.60(4H, m) 1.96(2H, m) 2.68(3H, s) 2.94(4H, m) 3.24(2H, dt, J=7.8, 6.3Hz) 3.80(3H, s) 5.05(1H, q, J=6.9Hz) 7.14-7.34(6H, m) 7.50(1H, s)
35			,
40	Compound	OH COOME	ONH COOMe
45	9	H Br	Br-(CH <sub>2</sub> ), Br-
50	Ex. No.	. Me	96 .
		<del> </del>	<del></del>

5	Elemental analysis (%)		
15	FAB-MS	441 (free base, MH <sup>†</sup> )	free base, MH <sup>+</sup> )
	IR (cm <sup>-1</sup> )		
	, 300MHz	12) 12) 5H2)	(2)
25	H-NMR & (ppm), 300MHz	CDCl <sub>3</sub> 1.45(2H, m) 1.55(2H, m) 1.55(2H, m) 2.46(3H, s) 2.67(3H, s) 2.94(2H, m) 2.94(2H, m) 3.19(1H, dd, J=14, 7Hz) 3.30(1H, dd, J=14, 5Hz) 3.00(1H, dd, J=8, 7, 5Hz) 6.74(1H, s) 7.16-7.34(5H, m) 7.83(1H, d, J=8Hz) 7.93(1H, s) 9.31(2H, brs) 11.84(1H, brs)	DMSO-d <sub>6</sub> 1.19(3H, t, J=7Hz) 1.30-1.63(6H, m) 2.85-2.90(5H, m) 3.17(1H, dd, J=14, 11Hz) 3.21(1H, dd, J=14, 5Hz) 4.14(2H, q, J=7Hz) 4.7(2H, q, J=16Hz) 4.88(1H, m) 7.17-7.34(5H, m) 8.13(1H, s) 8.60(2H, brs) 9.55(1H, d, J=8Hz) 13.14(1H, s)
30	H-N	CDC <sub>3</sub> 1.45(2H, m) 1.55(2H, m) 1.87(2H, m) 2.46(3H, s) 2.67(3H, s) 2.67(3H, s) 2.94(2H, m) 3.19(1H, dd, J 3.30(1H, dd, J 3.80(3H, s) 5.00(1H, ddd, G, 74(1H, s) 7.16-7.34(5H, s) 7.16-7.34(5H, s) 7.16-7.34(5H, s) 9.31(2H, brs) 11.84(1H, brs)	
35 .		/ Ph	- Ph - C00CH, C00Bt
40	Compound	OH CONH	CONH CONH
45		MeN-(CH <sub>2</sub> ), He	MeN-(CH <sub>2</sub> ), C1
50	Bx.	Men H H	- HCI

1		T	
5	Elemental analysis (%)		
	1		
15	Si	(free base, MH <sup>+</sup> )	ffee base, MH <sup>†</sup> )
20	IR (cm <sup>-1</sup> )		
20			
30 Table 52	<sup>1</sup> H-NMR & (ppm), 300MHz	CDCl <sub>3</sub> 1.38(3H, t, J=6.9Hz) 1.40-2.0(6H, m) 2.65(3H, s) 2.80-3.00(4H, m) 3.38(2H, d, J=6.3Hz) 4.37(2H, q, J=7.2Hz) 7.08(2H, d, J=8.7Hz) 7.08(2H, d, J=8.7Hz) 7.23-7.38(7H, m) 8.07(2H, d, J=8.9Hz) 9.44(2H, brs)	DMSO-d <sub>6</sub> 1.13(3H, t, J=9Hz) 1.20-1.60(7H, m) 1.80-1.87(2H, m) 2.70(3H, s) 2.80-2.95(4H, m) 3.10-3.40(4H, m) 4.11(2H, q, J=9Hz) 4.70(1H, m) 7.18-7.30(5H, m) 8.11(1H, s) 9.75(1H, brs) 13.15(1H, brs)
		CDCI <sub>3</sub> 1.38(3) 1.40-2 2.65(3) 2.80-3 3.38(2) 5.22(2) 7.08(2) 7.08(2) 7.08(2) 7.08(2)	DM 11.12 11.86 11.86 22.86 22.86 44.76 44.76 44.76 13.73
35		-Ocoost	COORT
		r <del>i</del> 000	CONH
40	Compound	HOONE HOO	13
			<b>_</b>
50		MeN-(CH;); H H CI	-HC1
	S.S.	. 66	8

5	Elemental analysis (%)		
15	FAB-MS	507 (free base, MH <sup>+</sup> ) (free base, MH <sup>+</sup> )	
	IR (cm <sup>-1</sup> )	KBr 3396 2933 2656	1734 1644 1589 1543 1405 1372 1254 1214 1099
25	<sup>1</sup> H-NMR & (ppm), 300MHz	Hz) (1) (2) (3) (4) (4) (4) (5) (4) (7) (7) (7) (8) (8) (9) (9) (9) (9) (9) (9) (9) (9) (9) (9	(Hz) 6.0, 7.2, 9.3Hz) 1.2Hz)
s Table 53	<sup>1</sup> H-NMR & (1	DMSO-d <sub>6</sub> 1.15(3H, t, 1=7.0Hz) 1.20-1.40(4H, m) 1.45-1.65(3H, m) 1.73-1.85(2H, m) 2.51(2H, s) 2.70-2.90(2H, m) 3.10-3.30(4H, m) 4.11(2H, q, 1=7.0Hz) 4.72(1H, dd, 1=6.0, 14, 7.15-7.30(5H, m) 8.13(1H, s) 8.87(1H, brs) 9.49(1H, brd, 1=7.1Hz) 1.12(3H, t, 1=7.1Hz) 1.95(2H, brs) 2.92(2H, t, 1=7.5Hz) 3.09-3.78(12H, m)	4.10(2H, q, J=7.1Hz) 4.71(1H, ddd, J=6.0, 7.2, 9.3Hz) 7.17-7.28(5H, m) 8.16(1H, s) 9.51(1H, d, J=7.2Hz) 9.64(2H, brs) 11.79(1H, brs) 13.20(1H, brs)
<b>35</b>		COOB!	
40	Compound	OH OH CONH	
45		(CH)	·2HC1 C1
50	Ex. No.	101 101 101	102

5	Elemental analysis (%)		
	FAB-MS	522.1 (free base, MH <sup>+</sup> )	439 (free base, MH <sup>+</sup> )
15	IR (cm <sup>-1</sup> )	9.	9
20			
Table 54	<sup>1</sup> H-NMR & (ppm), 300MHz	Ph DMSO-d <sub>6</sub> 1.13(3H, t, J=7.2Hz) 2.00Bt 1.77-2.12(6H, m) 2.85-3.27(7H, m) 3.48(1H, brd, J=8.6Hz) 4.10(2H, q, J=7.2Hz) 4.67-4.78(1H, m) 7.19-7.28(5H, m) 8.15(1H, s) 8.22(3H, brs) 10.27(1H, brs) 13.22(1H, brs)	CDCl <sub>3</sub> 0.90-2.05(20H, m) 2.50-2.70(5H, m) 2.96(2H, m) 3.82(1H, s) 4.82(1H, m) 6.68(1H, s) 7.45(1H, s)
30	H <sub>1</sub>	-Ph DMSO-d <sub>6</sub> 1.13(3H, t, 1.13(3H, t, 1.77-2.12(6) 2.85-3.27(7) 3.48(1H, b) 4.10(2H, g) 4.67-4.78(1) 7.19-7.28(2) 8.15(1H, b) 9.49(1H, b) 13.22(1H, t)	CDCl <sub>3</sub> 0.90-2.05(20 2.50-2.70(5H 2.96(2H, m) 3.82(1H, s) 4.82(1H, m) 6.68(1H, s) 7.45(1H, s)
35	Pu	OH CONH	Coogt
40	Compound	CH4), CH4)	OH CONH
45		11 N-(C	MeN-(CH2) 6 —— H C1/
50	No.	E01	20 M

89

5	tal ysis (%)	:	
	Elemental analysis		·
10	PAB-MS	568 (free base, MH*)	511 (free base. MH*)
15	IR (cm <sup>-1</sup> )		
Table 55	1H-NMR & (ppm), 300MHz	) J=8. 5Hz) 4H, m) 1H, m) 1H, m) 5H, m) 5H, m) J=7. 6Hz) J=7. 6Hz)	-d, 381, 8) 341, 8) 341, 8) 342, 8) 343, 10(41, m) 343, 8) 344, 8) 344, 8) 347
25 <b>E</b>	1H-NMR & (p	DMSO-de 2. 52(3H, b) 2. 52(3H, d) 2. 86-3. 02(3, d) 3. 48(3H, s) 4. 09(2H, b) 4. 14-7, 32(7, 14-7, 32(7, d) 7. 56-7. 68(7, 72-7, 82(8, d) 9. 04(1H, d)	DMSO-de 1.84(4H, s) 2.17(3H, s) 2.55(2H, s) 2.95-3.10(4H, m) 3.40(3H, s) 4.05(2H, s) 4.05(2H, s) 7.20-7.35(5H, m) 7.50(1H, s) 8.66(1H, brs) 8.91(1H, d. J=9.0
30		Ph	Ph COOMe
35	Compound	THE TOP OF	CONTE
40	0	C1. H C1.	C1 CH3)4-0-(CH3)4-0-(C1)-(C1)-(C1)-(C1)-(C1)-(C1)-(C1)-(C1)
45			
	Bx. No.	105	106

5	Elemental analysis (%)	CreHratlaNrO.+HCl Calculated C. 54, 22 H. 5. 78 N. 4. 66 Pound C. 54, 24 H. 5. 75 N. 4. 83	
15	FAB-MS	539 (free base, MH+)	553 (free base, MH+)
. 20	IR (cm <sup>-1</sup> )	KBr 3285 2850 2723 1768 1745 1648	Neat 2957 1749 1666 1456
72 Table 56	1 H-NMR & (ppm), 300MHz	DMSO-d <sub>4</sub> 15(3H, d, J=6. 0Hz) 17(3H, d, J=6. 0Hz) 40-1. 90(4H, m) 50-2. 58(3H, m) 65-2. 73(1H, m) 90-3. 17(3H, m) 63(3H, s) 63(3H, s) 63(3H, s) 70-4. 65(2H, m) 55-4. 62(1H, m) 55-4. 62(1H, m) 70-8. 85(2H, m) 70-8. 85(2H, m) 95(1H, d, J=7. 0Hz)	DMSO-d <sub>6</sub> (400MHz) 1. 22(9H, s) 1. 78-1. 90(4H, m) 2. 53(3H, m) 2. 90-3. 03(3H, m) 3. 13(1H, dd, J=13. 82, 5. 53Hz) 3. 62(3H, s) 4. 00-4. 08(2H, m) 4. 53-4. 60(1H, m) 7. 20-7. 32(5H, m) 7. 38(1H, s) 8. 82(2H, hrs) 8. 82(2H, hrs) 8. 97(1H, d, J=7. 80Hz)
35		CONH Ph COOKe 1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.	CONH COOME 3.1.1.2.9.1.1.2.9.1.1.2.9.1.1.2.9.1.1.1.2.9.1.1.1.2.9.1.1.2.9.1.1.2.9.1.1.2.9.1.1.2.9.1.1.2.9.1.1.2.9.1.1.2.9.1.1.2.9.1.1.2.9.1.1.2.9.1.1.2.9.1.1.2.9.1.1.2.9.1.2.2.9.1.2.2.9.1.2.2.2.2
40	Compound		13 0 13
45		MeN-(CH <sub>s</sub> ), H +HC1	MeN-(CH2)4 H •HC1
50 .	Š.	107	108

E
J

	Elemental analysis (%)		
•	FAB-MS	540 (free base, MH+)	571 (free base, M*H)
	IR (cm <sup>-1</sup> )	KBr 3422 2954 1741 1646 1456	
Table 57	1H-NNR & (ppm), 300MHz	DNSO-d <sub>4</sub> 1. 90-1. 97(4H, m) 2. 53(3H, t, J=6Hz) 2. 99-3. 12(7H, m) 3. 16(1H, dd, J=12. 6Hz) 3. 66(3H, s) 4. 00-4. 10(2H, m) 4. 56-4. 65(1H, m) 7. 20-7. 34(5H, m) 7. 20-7. 34(5H, m) 7. 57(1H, s) 8. 12(3H, s) 8. 12(3H, s) 9. 08(1H, d, J=6Hz)	DMSO-d <sub>4</sub> 1. 84(4H, bs) 2. 11(3H, s) 2. 49(2H, bs) 2. 49(2H, bs) 2. 88-3. 22(4H, m) 3. 63(3H, s) 4. 65(2H, bs) 4. 52-4. 68(1H, m) 7. 12-7. 34(5H, m) 7. 54(1H, s) 8. 85(2H, bs) 8. 99(1H, d, J=7. 6Hz)
	Compound	Men-(CH <sub>2</sub> ) <sub>4</sub> -0 - CONH - COOMe H C1	C1 0 -CH <sub>2</sub> OAC Ph MeN-(CH <sub>2</sub> ) <sub>2</sub> -0 -CONH -C00Me H c1
	S	109	110

5	Blemental analysis (%)		C2,44,6C12N2O4.HC1 Calculated C,56.55 H, 6.05 N, 4.55 Round C,56.17 H, 6.16 N, 4.48
	FAB-MS	659 (free base, MH+)	579 (free base, MH+)
20	IR (cm <sup>-1</sup> )	Neat 2954 2728 1778 1739 1667	KBr 3422 2935 1745 1654 1452
Table 58	1 H-NMR & (ppm), 300MHz	MSO-d <sub>6</sub> 89-1. 96(4H, m) 54(3H, brs) 65-2. 82(4H, m) 90-3. 05(3H, m) 14(1H, dd, J=15, 3Hz) 62(3H, s) 62(3H, s) 12(2H, s) 12(2H, s) 18-7. 40(10H, m) 50(1H, s) 77(2H, brs) 94(1H, d, J=9Hz)	16, 17, 17, 17, 17, 17, 17, 17, 17, 17, 17
30	1 H-NMR	DMSO-de 1. 89-1. 96(4) 2. 54(3H, brs 2. 65-2. 82(4) 2. 90-3. 05(3) 3. 14(1H, dd, 3) 4. 50-4. 65(1) 5. 12(2H, s) 7. 18-7. 40(1) 7. 18-7. 40(1) 7. 50(1H, s) 8. 77(2H, brs 8. 94(1H, d, J)	DMSO-de 1. 1-1. 9(14H 2. 49-2. 51(1 2. 54(3H, s) 2. 93-3. 17(4 3. 63(3H, s) 4. 0-4. 10(2H 4. 55-4. 15(1 7. 23-7. 32(5 7. 44(1H, s) 8. 95(1H, d, J
35		-(CH <sub>2</sub> ) <sub>2</sub> C00Bn	MH COOMe
40	Compound	0 10	C1 CONH
45		MeN-(CH2),4-0 H •HC1	MeN-(CH2),-0 H •HCl
50	Bx. No.	H	112

5		Elemental analysis (%)		C3.0H3.2Cl2N2O6.HCl Calculated C, 57.75 H, 5.33 N, 4.49 Found C, 57.71 H, 5.31 N, 4.47
15		FAB-MS	587 (free base, MH+)	587 (free base, MH+)
20		1R . (cm <sup>-1</sup> )	Neat 2953 1747 1663 1453	KBr 3433 2948 2719 1744 1645 1457
25 30	Table 59	1H-NMR & (ppm), 300MHz	DMSO-d <sub>6</sub> 1. 78-1. 96(4H, m) 2. 75(6H, brs) 2. 95(1H, dd, J=15, 9Hz) 3. 05-3. 16(3H, m) 3. 48(3H, s) 4. 10(2H, t, J=6Hz) 4. 48-4. 56(1H, m) 7. 17-7. 29(5H, m) 7. 17-7. 29(5H, m) 7. 73-7. 81(1H, m) 8. 00-8. 05(2H, m) 9. 04(1H, d, J=6Hz) 10. 05(1H, brs)	DMSO-d <sub>4</sub> 1. 8-1. 9(4H, m) 2. 53(3H, s) 2. 80-3. 15(4H, m) 3. 32(3H, s) 3. 32(3H, s) 4. 05-4. 10(2H, m) 4. 51-4. 60(1H, m) 7. 19-7. 61(9H, m)
35		Ħ <sub>1</sub>	-Ph -C00Me -C00Me -2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.	-Ph 22. 22C00Me 3. 4. 4. 4. 7. 7. 7.
40		Compound	CI CONTH	O We CONH
45			Me2N-(CH2),4-C	CI MeN-(CH*),-0 H CI
50		Bx. No.	113	114

5		Elemental analysis (%)	Cs. 243, Cl. 2N2 04. + HCl Calculated C, 58. 95 H, 5. 72 N, 4. 30 C, 58. 95 H, 5. 98 N, 4. 21	
15		FAB-MS	615 (free base, NH*)	541 (free base, MH+)
20		IR (cm <sup>-1</sup> )	KBr 1748 1455 1211 1057	KBr 3423 2955 1774 1746 1669 1215 1029
25	09 e	pm), 300MHz	l. m) l. m) l. m) =13. 5, 6. 0Hz) l. m) l. m) l. m)	J=7. 6H2) J=5. 6H2) 4H, m) J=7. 6H2) IH, m) 5H, m) 5H, m)
30	Table 60	1H-NMR & (ppm), 300MHz	DMSO-de 1. 76-1. 94(4H, m) 2. 23(3H, s) 2. 23(3H, s) 2. 30(6H, s) 2. 50-2. 58(3H, m) 2. 86-3. 03(3H, m) 3. 11(1H, dd, J=13. 5, d) 4. 02-4. 12(2H, m) 4. 52-4. 63(1H, m) 6. 98(2H, s) 7. 13-7. 31(5H, m) 7. 13-7. 31(5H, m) 7. 41(1H, s) 8. 69(1H, brs) 9. 08(1H, d, J=6. 0Hz)	DMSO-de 1. 26(3H, t, J=7, 1. 85(4H, bs) 2. 85(3H, t, J=5, 2. 84-3, 22(4H, 1, 3. 64(3H, s) 4. 22(2H, q, J=7, 4. 22(2H, q, J=7, 7. 20-7, 38(5H, 1, 7. 50-7, 38(5H, 1, 8. 76(2H, brs) 9. 01(1H, d, J=8,
35			- Ne - COOMe	/ Ph
40	·	Compound	O We CONH	O CONH
45	·	0	C1 MeN-(CH <sub>2</sub> ),-0-1 H C1	C1)  H  C1  -HC1
50		No.		116 Me

5	Blemental analysis (%)	C2sHsoCl2N2O4·HCl Calculated C, 53, 44 H, 5, 56 N, 4, 99 Pound C, 52, 79 H, 5, 46 N, 4, 94	C2,442,C12N2O4.HC1 Calculated C, 54, 22 H, 5, 78 N, 4, 86 Found C, 54, 04 H, 5, 68 N, 5, 01
15	PAB-MS	525 (free base, MH <sup>+</sup> )	539 (free base, MH*)
20	IR (cm <sup>-1</sup> )	KBr 1646 1528 1456 1372 1190	KBr 1734 1655 1456 1373 1201
rable 61	'H-NMR & (ppm), 300MHz	DMSO-d <sub>4</sub> 1. 15(3H, t, J=6. 0Hz) 1. 76-1. 91(4H, m) 2. 17(3H, s) 2. 50-2. 59(3H, m) 2. 90-3. 07(3H, m) 3. 14(1H, dd, J=6. 0, 15. 0Hz) 4. 00-4. 15(4H, m) 4. 55-4. 62(1H, m) 7. 20-7. 33(5H, m) 7. 20-7. 33(5H, m) 7. 51(11H, s) 8. 69(2H, brs) 8. 89(1H, d, J=9. 0Hz)	90-de 9-1. 12(3H, d, J=6. 0Hz) 9-1. 12(3H, d, J=6. 0Hz) 1-1. 92(4H, m) 1-2. 92(4H, m) 1-2. 59(3H, brs) 1-3. 17(3H, m) 1-4. 17(3H, m) 1-4. 12(2H, m) 1-4. 12(2H, m) 1-4. 60(1H, m) 1-7. 36(5H, m) 1-7. 36(5H, m) 1-1. 8) 1(1H, d. J=9. 0Hz) 1(1H, d. J=9. 0Hz)
30 Eg	H-NMR	DMSO-d <sub>6</sub> 1. 15(3H, t) 1. 76-1. 91 2. 17(3H, s) 2. 17(3H, s) 2. 90-2. 59 2. 90-2. 59 3. 14(1H, d) 4. 65-4. 62 4. 55-4. 62 7. 20-7. 33 7. 51(1H, s) 8. 69(2H, s) 8. 89(1H, d)	DNSO-de 0. 95-1. 0 1. 78-1. 9 2. 18(3H, 9 2. 51-2. 5 2. 39-3. 1 4. 48-4. 0 4. 88(1H, 6 7. 19-7. 3 7. 52(1H, 8 8. 77(1H, 8
<b>35</b>		Ph coogt	# C00
40	Compound		
<b>45</b>		C1, MeN-(CH <sub>2</sub> ),-0 H C1'	C1. MeN-(CH <sub>2</sub> ),-0 H .HC1
50	S. S.	711	118

5	Elemental analysis (%)	C1.0H1.Cl1.N2O4.HCl Calculated C. 57. 75 H. 5. 33 N. 4. 49 Pound C. 56. 70 H. 5. 21 N. 4. 36	
15	FAB-MS	539 (free base. MH+)	601 (free base, MH+)
20	IR (cm <sup>-1</sup> )	KBr 3420 2980 1749 1669 1522 1452	Neat 2980 1746 1668 1453
25 Zapie 62 30	H-NMR & (ppm), 300MHz	DMSO-d <sub>6</sub> 1. 00(3H, d, J=6Hz) 1. 04(3H, d, J=6Hz) 1. 75-1. 90(4H, m) 2. 82-3. 09(4H, m) 4. 05-4. 12(2H, m) 4. 40-4. 50(1H, m) 4. 70-4. 85(1H, m) 7. 05-8. 05(14H, m) 9. 02(1H, d, J=7. 0Hz)	DMSO-d <sub>4</sub> 1. 00(3H, d. J=6Hz) 1. 04(3H, d. J=6Hz) 1. 80-1. 93(4H, m) 2. 54(3H, t. J=6Hz) 2. 54(3H, t. J=6Hz) 2. 90-3. 00(3H, m) 3. 05(1H, dd, J=15, 6Hz) 4. 05-4. 13(2H, m) 4. 40-4. 50(1H, m) 4. 74-4. 81(1H, m) 7. 17-7. 29(5H, m) 7. 17-7. 29(5H, m) 7. 17-7. 29(5H, m) 8. 00-8. 05(2H, m) 8. 00-8. 05(2H, m) 9. 02(1H, d. J=9Hz)
35		Ha Coo	Ph C00
40	Compound	No contraction	CONI
45		C1 H2N-(CH2)4-0- C1	C1 MeN-(CH <sub>2</sub> ),-0- H C1
50	S. S.	1 61	021

45	Co	C1 CHs) 2-0 — C1	C1 HN N-(CH <sub>2</sub> ),-0 -
40	Compound	CONTI	CONN
. 35		DMSO-de 1. 28(3H, d. J=6. 40Hz) 2. 17(3H, s) 2. 90-3. 90(11H, m) 4. 38-4. 40(2H, m) 4. 45-4. 61(1H, m) 7. 15-7. 30(5H, m) 7. 52(1H, s) 7. 52(1H, s) 7. 55(1H, d. J=7. 0Hz) 9. 50-9. 80(2H, m)	COOM®
Table 63	1H-NARS (p	DMSO-de 1.28(3H, d, J= 2.17(3H, s) 2.90-3.90(11 4.88-4.40(2H 4.45-4.61(1H 7.15-7.30(5H 7.75(1H, d, J= 9.50-9.80(2H	DMSO-de 1. 29(3H, S) 2. 18(1H, S) 3. 02(1H, d, J= 3. 02(1H, d, J= 3. 05(3H, S) 3. 65(3H, S) 4. 55-4. 68(1H 4. 55-4. 68(1H 7. 18-7. 37(5H 7. 52(1H, S) 9. 58(2H, d, J= 9. 58(2H, brs)
25	1 H-NMR & (ppm), 300MHz	6. 40Hz) H. m) I. m) I. m) I. m) T. 0Hz)	DMSO-de 1. 29(3H, d. J=6. 3Hz) 2. 18(1H, s) 3. 02(1H, d. J=15. 0, 8. 5Hz) 3. 02(1H, d. J=15. 0, 6. 0Hz) 2. 90-3. 80(9H, m) 3. 65(3H, s) 4. 55-4. 68(1H, m) 7. 18-7. 37(5H, m) 7. 52(1H, s) 8. 92(1H, d. J=14. 0Hz) 9. 58(2H, brs)
20	IR (cm <sup>-1</sup> )		KB7 3422 1742 1664 1455 1188 1118
15	FAB-MS	537 (free base, MH*)	551 (free base. MH*)
5 10	Elemental analysis (%)		Ca. Ha. Clana 0 . 2 HCl Calculated C. 49. 94 H. 5. 32 N. 6. 72 Pound C. 48. 39 H. 5. 16 N. 6. 46

5	Elemental analysis	Optical rotation: [α] <sup>3*</sup> <sub>b</sub> = -26.8° (c=1.01, MeOH)	Cai HaaClanaOa.2HCl Calculated C. 49. 42 H. 5. 36 N. 7. 20 Found C. 48. 47 H. 5. 58 N. 6. 91
15	PAB-MS	614 (free base, MH+)	614 (free base, MH+)
20	(CM-1)	KBr 3430 1747 1664	KBr 3422 1741 1642 1585 1458 1357 1221
rable 64	'H-NMR & (ppm), 300MHz	DMSO-d <sub>6</sub> 1. 29(3H, d, J=6. 3Hz) 2. 95(1H, dd, J=9. 9, 13. 8Hz) 3. 09(1H, dd, J=9. 9, 13. 8Hz) 3. 20-3. 80(8H, m) 3. 48(3H, s) 4. 49-4. 55(2H, m) 7. 18-7. 28(5H, m) 7. 56(1H, s) 7. 62(2H, t, J=7. 8Hz) 7. 78(1H, t, J=7. 5Hz) 7. 53(2H, d, J=8. 4Hz) 9. 06(1H, d, J=7. 8Hz)	DMSO-d. 1. 29(3H, d. J=6. 2Hz) 2. 95(1H, dd. J=13. 8. 9. 8Hz) 3. 09(1H, dd. J=13. 8. 5. 4Hz) 3. 12-3. 93(9H, m) 3. 48(3H, s) 4. 42-4. 57(3H, m) 7. 18-7. 29(5H, m) 7. 56-7. 64(3H, m) 7. 75-7. 80(1H, m) 8. 01-8. 04(2H, m) 9. 05(1H, d. J=7. 8Hz) 9. 55-9. 87(2H, m)
35		-conh — coone	Ph Ph COOMe
40	Compound		5 0 5
45		HN N- (CH2)3- Me N- (CH2)3 2HC1	HN N-(CH <sub>2</sub> ) <sub>2</sub> -
50	Bx.	123 M	124 M

Table 65

1						
8.8.	Compound	<sup>1</sup> H-NMR & (ppm), 300MHz	(cm <sup>-1</sup> )	FAB-MS	Elemental analysis (%)	
83	Me-N N-(CH <sub>2</sub> ) <sub>2</sub> -0 CONH C00Et	DMS0-d <sub>4</sub> 1. 15(3H, t, J=7. 3Hz) 2. 18(3H, s) 2. 80(3H, s) 3. 02(1H, dd, J=9. 5, 13. 8Hz) 3. 15-3. 68(10H, m) 4. 09(2H, q, J=7. 3Hz) 4. 38(2H, bs) 4. 59(1H, ddd, J=5. 7, 9. 59(1H, ddd, J=5. 7, 13. 8Hz) 7. 23-7. 33(5H, m) 7. 53(1H, s)	KBr 3433 2984 2418 1769 1735 1666 1529 1456 1195 1195	566 (free base, MH+)		<u> </u>
		8. 91(1H, d, J=7, 6Hz)				

5		Elemental analysis (%)		C1.0H1.2C11N1.05.HC1 Calculated C. 50. 27 H. 4. 85 N. 5. 86 Found C. 50. 22 H. 5. 16 N. 5. 47
15		FAB-MS	455 (free base, MH*)	441 (free base, 岷(*)
20		[R (cm <sup>-1</sup> )		KBr 2971 1638 1585 1541 1457 1221
25	99 a	¹H-NMR & (ppm), 300MHz	) J=5. 6Hz) 2H. m) 1H. m) 5H. m) J=7. 7Hz)	(4H, m) (2H, m) (2H, m) (1, J=15, 0, 10, 5Hz) (2H, m) (1H, m) (5H, m) (5H, m) (5H, m) (5H, m) (5) (5)
30	Table 66	1H-NMR & (p	DNSO-d. 1. 83(4H, bs) 2. 54(3H, t. J=5. 6 2. 95(2H, bs) 3. 08-3. 78(2H, m) 4. 05(2H, bs) 4. 62-4. 72(1H, m) 7. 20-7. 38(5H, m) 8. 21(1H, s) 8. 21(1H, s) 9. 36(1H, d, J=7. 7 13. 47(1H, s)	DMSO-4. 1. 69-1. 92(4H) 2. 79-2. 96(2H) 3. 09(1H, dd. J) 3. 26(1H, dd. J) 3. 96-4. 11(2H) 4. 63-4. 77(1H) 7. 13-7. 35(5H) 7. 86(3H, brs) 8. 20(1H, s) 13. 46(1H, s)
35			4 C00H	Ph C00H
40		Compound	HO CONH	HO CONH
45	• .	වී	CI CI	C1 C1 C1
50		Bx. No.	MeN-( H 126 ·HC1	H <sub>2</sub> N

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Table 67

		10 01001			
S. S.	Сотроипа	'H-NAR & (ppm), 300AHz	IR (cm <sup>-1</sup> )	FAB-MS	Elemental analysis (%)
128	C1 OH Ph Me <sub>3</sub> N-(CH <sub>2</sub> ) <sub>4</sub> -0 -C0NH -C00H OHC1	DMSO-d <sub>6</sub> 1. 70-1. 90(4H. m) 2. 76(6H. s.) 3. 10-3. 40(4H. m) 4. 04-4. 08(2H. t, J=7Hz) 4. 60-4. 75(1H. m) 7. 19-7. 30(5H. m) 8. 18(1H. s.) 9. 41(1H. brs.)	KBr 3422 1735 1638 1584 1458	(free base, MH+)	·
129	HN N-(CH <sub>2</sub> ) <sub>2</sub> -0 CONH COOH  Me  C1  C1  CH  COOH  -2HC1	CD <sub>3</sub> CO <sub>2</sub> D 1.51(3H, s) 3.10-3.40(2H, m) 3.70-4.30(9H, m) 4.51-4.60(2H, m) 5.09-5.06(2H, m) 7.29-7.21(5H, m) 7.94(1H, s)	KBr 3418 2941 1734 1641 1457	495 (free base, NH*)	CasHarClansOs-2HCl Calculated C. 48. 52 H. 5. 13 N. 7.38 Pound C. 47. 55 H. 5. 02 N. 6. 72

r	·		
5	Blemental analysis (%)		
15	FAB-MS	419 (free base, MH <sup>†</sup> )	(free base MH <sup>+</sup> )
20	IR (cm <sup>-1</sup> )	KBr 3368 2940 1733 1639 1543 1408 1258 1203 701	
25	<sup>1</sup> H-NMR & (ppm), 300MHz	DMSO-d <sub>6</sub> 1.28-1.38(2H, m) 1.50-1.64(4H, m) 2.50(3H, s) 2.62(2H, t, J=7.5Hz) 2.80-2.89(2H, m) 3.07(1H, dd, J=8.9, 13.9Hz) 3.20(1H, dd, J=4.8, 13.9Hz) 4.69(1H, dd, J=4.8, 7.8, 8.9Hz) 6.89(1H, s) 7.12-7.30(5H, m) 7.94(1H, s) 8.59(2H, brs) 8.98(1H, d, J=7.8Hz) 12.05(1H, brs) 12.05(1H, brs)	ଜିଜିଜିଜି ଜି
30 Table 68	<sup>1</sup> H-NMR	DMSO-d <sub>6</sub> 1.28-1.38(2H, m) 1.50-1.64(4H, m) 2.50(3H, s) 2.62(2H, t, J=7.5Hz) 2.80-2.89(2H, m) 3.07(1H, dd, J=8.9, 1) 3.20(1H, dd, J=4.8, 1) 4.69(1H, dd, J=4.8, 1) 7.12-7.30(5H, m) 7.94(1H, s) 8.59(2H, brs) 8.59(2H, brs) 12.05(1H, brs) 12.05(1H, brs)	DMSO-d <sub>6</sub> 1.25-1.40(2H, m) 1.50-1.70(4H, m) 2.80-2.70(8H, m) 2.94-3.40(4H, m) 4.68(1H, m) 6.90(1H, s) 7.09-7.20(5H, m) 7.95(1H, s) 8.97(1H, brs)
35		-Рћ -соон	Ph C000H
40	Compound	HOOD—	OH CONH
45		CH*)*	Me.N-(CH.), Cl
50	*0	MeN-( H +HC1	Me.N-
	RX.	71	<u> </u>

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5	Blemental analysis (%)		
. 15	FAB-MS	453 (free base, MH <sup>+</sup> )	467 (free. base, MH <sup>+</sup> )
	IR (cm <sup>-1</sup> )		
25 Q Q	<sup>1</sup> H-NMR δ (ppm), 300MHz	H, m) H=7.4Hz)	m) J=9, 14Hz) J=5, 14Hz) J=5, 8Hz) m)
30	H-NM	DMSO-d <sub>6</sub> 1.32-1.64(6H, m) 2.85(4H, m) 3.12-3.34(2H, m) 3.57(3H, s) 4.68-4.72(1H, m) 7.16-7.30(5H, m) 8.13(1H, s) 8.65(2H, brs) 9.38(1H, d, J=7.4Hz) 13.14(1H, brs)	DMSO-d <sub>6</sub> 1.35(2H, m) 1.45-1.6(4H, m) 2.59(6H, s) 2.75(2H, m) 2.83(2H, m) 2.83(2H, dd, J=9, 14H 3.13(1H, dd, J=5, 14H 4.62(1H, dd, J=5, 8Hz) 7.15-7.2(2H, m)
35		-Ph	- Ph
40	Compound	CONH CONH	OH CONH
45		C1) H C1 -HC1	.HCI
50	Bx. No.	MeN- H 132 •HC	Me <sub>2</sub>
	<u> </u>	<del></del>	L

	Bx.	134	135
50		Me-NN-(CH <sub>2</sub> ) <sub>2</sub> -	He-l
45	3	C1.	CI.
	Compound	OH CONH	CI
35		Ph C003	H COOH
30 E	1H-NMR	DMSO-d <sub>6</sub> 2.79(3H, s) 3.04-4.10(12H, m) 4.33(2H, brs) 4.70(1H, m) 7.17-7.30(5H, m) 8.21(1H, s) 9.36(1H, d, J=8Hz) 13.47(1H, brs)	
25	H-NMR & (ppm), 300MHz	H, m) (m, t) (m, t) (m, t) (m, t) (1) (1) (1) (2) (3)	DMSO-4 <sub>6</sub> 1.36(2H, m) 1.69(2H, dd, 1=6.3, 12.6Hz) 1.75-2.0(3H, m) 2.67(3H, 8) 2.87(2H, m) 2.95(1H, dd, 1=8.4, 14.5Hz) 3.11(1H, dd, 1=5.4, 14.1Hz) 3.93(2H, t, 1=6.3Hz) 7.17-7.29(5H, m) 7.59(1H, s)
20	IR (cm <sup>-1</sup> )		
15		496 (free base, MH <sup>+</sup> )	(free base, MH <sup>+</sup> )
5	FAB-MS Elemental analysis (%)		
	_		

5	
10	
15	
20	
25	
30	

7
<u>=</u>
Table
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				-	
Bx.	Compound	'H-NMR & (ppm), 300MHz	IR (cm⁻¹)	PAB-MS	Elemental analysis (%)
13%	Men-(CH <sub>2</sub> ),-N CONH COOMe H C1 -2HC1	DMSO-d <sub>4</sub> 1. 44-1. 68(4H, m) 2. 50-2. 57(3H, m) 2. 63-2. 92(2H, m) 3. 11(1H, dd. J=13. 5, 9. 0Hz) 3. 21(1H, dd. J=13. 5, 6. 6Hz) 3. 48-2. 59(2H, m) 3. 05(3H, s) 4. 67-4. 79(1H, m) 5. 63(1H, brs) 7. 17-7. 34(5H, m) 8. 02(1H, s) 8. 57(2H, brs) 9. 18(1H, d. J=9. 0Hz) 13. 43(1H, s)		468 (free base, MH+)	
137	MeN-(CH <sub>2</sub> ) <sub>6</sub> -N — CONH — COOMe H + H + COOMe	DMSO-d <sub>4</sub> 1. 29-1. 84(8H, m) 2. 69(3H, s) 2. 88-3. 36(6H, m) 3. 73(3H, s) 4. 82-4. 96(1H, m) 6. 93(1H, d, J=8. 5Hz) 7. 18-7. 34(6H, m) 7. 93(1H, t, J=4. 2Hz)	KBr 3422 2939 1741 1638 1542	428 (free base, M⁺H)	

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5	Elemental analysis (%)		C2.0H2.8C1.2N5.04.HC1 Calculated C.50.38 H. 5.07 N. 8.81 Pound C. 47.87 H. 4.6 N. 7.31
10			00 - 2 - 2
15	FAB-MS	516 (free base, MH*)	440 (free base, MH+)
20	[R (cm <sup>-1</sup> )	KBr 3412 2954 1638 1599 1542 1445 1066	KBr 2955 1677 1458 1413 1352 1261 1203 1138
25 E	H-NMR & (ppm), 300MHz	DMSO-de 1. 83(4H, bs) 2. 82-2. 94(2H, m) 3. 16-3. 32(2H, m) 4. 02-4. 06(2H, m) 4. 88-5. 02(1H, m) 7. 06-7. 42(8H, m) 7. 62(2H, d, J=8. 1Hz) 7. 86(2H, bs) 8. 32(1H, s) 10. 38(1H, s)	DMSO-ds 1. 74-1. 87(4H, m) 2. 83-2. 92(2H, m) 2. 95-3. 03(1H, m) 3. 14-3. 22(1H, m) 3. 99-4. 06(2H, m) 4. 65-4. 74(1H, m) 7. 14-7. 34(6H, m) 7. 68-7. 81(4H, m) 8. 23(1H, s) 9. 19-9. 21(1H, m) 13. 56(1H, s)
35		NH CONH-Ph	INH CONH2
40	Compound	# J3	CI
45		H <sub>2</sub> N-(CH <sub>2</sub> ),-0	H <sub>2</sub> N-(CH <sub>2</sub> ),-0
50	RX.	138	681

5	Elemental analysis (%)	Ca. Has Clans 04 - HCl Calculated C, 51. 39 H, 5. 34 N, 8. 56 Pound C, 50. 03 H, 5. 38 N, 8. 15	
15	PAB-MS	454 (free base. MH <sup>+</sup> )	534 (free base, MH+)
20	IR (cm <sup>-1</sup> )	KBr 3422 2940 2940 1641 1412 1348 1228	KBr 2954 1670 1639 1542 1508 1217 1065
73 table 73	1H-NMR & (ppm), 300MHz	DMSO-d <sub>6</sub> 1. 82 (3H, d. J=4. 5Hz) 2. 62 (3H, d. J=4. 5Hz) 2. 85-2. 89 (2H, m) 3. 00 (1H, dd, 3. 16 (1H, dd, 1=13. 7, 4. 2Hz) 4. 00-4. 16 (2H, m) 7. 13-7. 32 (5H, m) 8. 20 (1H, q. J=4. 5Hz) 8. 27 (1H, s) 9. 31 (1H, d. J=8. 2Hz) 13. 56 (1H, s)	DMSO-d <sub>4</sub> 1. 78-1. 86(4H, m) 2. 78-2. 94(2H, m) 3. 18-3. 78(2H, m) 4. 02-4. 10(2H, m) 7. 12-7. 42(8H, m) 7. 12-7. 42(8H, m) 7. 32(2H, bs) 8. 32(1H, s) 9. 42(1H, d, J=8. 8Hz) 10. 44(1H, s)
30			Ľ.
35	pur	Ph CONFI-Me	Ph CONFI
40	Compound	HO OH	HO
45		C1 H2N-(CH2),-0 C1	C1 H2N-(CH2),-0 C1
50	Bx. No.	140	H 141

10 .	IS Blemental analysis (%)	CasHaeClaN404.HCl Calculated C. 50.87 H. 4. 78 N. 9.49 Found C. 49.81 H. 5.14 N. 9.27	
15	FAB-MS	517 (free base. MH*)	470 (MH+)
20	[R (cm <sup>-1</sup> )	KBr 3423 2957 2957 1543 1541 1439 1260 1228	KBr 3422 1624 1570 1542 1431
Table 74	H-NAR S (ppm), 300AHz	DMSO-d <sub>4</sub> 1. 73-1. 88(4H, m) 2. 79-2. 92(2H, m) 3. 08-3. 30(2H, m) 4. 01-4. 07(2H, m) 5. 02-5. 32(1H, m) 7. 17-7. 21(2H, m) 7. 26-7. 31(2H, m) 7. 45-7. 47(2H, m) 7. 45-7. 47(2H, m) 7. 45-7. 47(2H, m) 7. 84-8. 08(5H, m) 8. 30(1H, s) 8. 37(1H, d, J=6. 0Hz) 11. 24(1H, s) 13. 42(1H, s)	DMS0-d <sub>6</sub> 1. 70-1. 90(4H, m) 2. 58(3H, s) 2. 75-3. 00(4H, m) 3. 88-3. 98(2H, m) 4. 56-4. 59(1H, m) 7. 16-7. 33(5H, m) 7. 55(1H, s) 8. 78(1H, m)
35		-Ph -CONH N	- CONHOH
40	Compound	OH CONH	T OH CONH
45		C1, H <sub>2</sub> N-(CH <sub>2</sub> ), -0 — C1'	C1.) MeN-(CH <sub>2</sub> ),-0 — H
50	Š. Š.	142	143

5		Elemental analysis (%)	
15		FAB-MS	479 (free base, MH+)
20		IR (cm <sup>-1</sup> )	KBr 3421 2935 1638 1588 1542 1457
<i>25</i>	Table 75	'H-NMR & (ppm), 300MHz	DMSO-d <sub>4</sub> 1. 80-1. 86(4H, m) 2. 35(3H, s) 2. 84-2. 90(2H, m) 3. 34-3. 47(2H, m) 4. 00-4. 06(2H, m) 5. 56-5. 64(1H, m) 7. 19-7. 34(5H, m) 7. 96(3H, brs) 9. 86(1H, d, J=9Hz) 13. 19(7H, brs)
35			Ph N N N N N N N N N N N N N N N N N N N
40		Compound	CI OH CONH
<b>45</b>			H <sub>2</sub> N-(CH <sub>2</sub> ),-0
50		33	

Š.	Compound	'H-NAR & (ppm), 300MHz	[R (cm <sup>-1</sup> )	FAB-MS	Elemental analysi
144	H <sub>2</sub> N-(CH <sub>2</sub> ) <sub>4</sub> -0 CONH N-(CH <sub>2</sub> ) <sub>4</sub> -0 CONH N-(CH <sub>2</sub> ) <sub>4</sub> -0 CONH N-(CH <sub>2</sub> ) <sub>1</sub> N-(CH <sub>2</sub> ) <sub>1</sub> N-(CH <sub>2</sub> ) <sub>1</sub> N-(CH <sub>2</sub> ) <sub>2</sub> N-(CH <sub></sub>	DMSO-d <sub>4</sub> 1. 80-1. 86(4H, m) 2. 35(3H, s) 2. 35(3H, s) 2. 84-2. 90(2H, m) 3. 34-3. 47(2H, m) 4. 00-4. 06(2H, m) 5. 56-5. 64(1H, m) 7. 19-7. 34(5H, m) 7. 19-7. 34(5H, m) 8. 22(1H, s) 9. 86(1H, d, J=9Hz) 13. 19(1H, brs)	KBr 3421 2935 1638 1588 1542 1457	479 (free base, MH+)	
145	C1 OH CH2)4-0-CH2 OH CH2 OH	DMSO-de 1. 70-1. 92 (4H, m) 2. 73-3. 01 (4H, m) 3. 42-3. 58 (2H, m) 3. 95-4. 11 (2H, m) 4. 13-4. 32 (1H, m) 4. 97 (1H, brs.) 7. 09-7. 33 (5H, m) 7. 91 (3H, brs.) 8. 25 (1H, s.) 8. 92 (1H, d. J=9. 0Hz.) 13. 98 (1H, s.)	KBr 3421 2950 1637 1583 1458	427 (free base, MH+)	Ca. M. 4 Cl 2 N. 2 O. 4 · HCl Calculated C. 51. 80 H. 5. 43 N. 6. 04 Pound C. 50. 96 H. 5. 46 N. 5. 65

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5		Elemental analysis (		
10		FAB-MS	437 (free base, MH+)	475 (free base, MH+)
		IR (cm <sup>-1</sup> )	KBr 3386 2952 1741 1647 1618 1527 1227 1188	KBr 3332 2723 2723 1750 1630 1605 11535 1204 1183
20		), 300MHz	. m)	J-5. 4Hz) 4H; a) 3H; a) 2H; a) 2H; a) 2H; a) 1-4. 3Hz) 1H; a)
25	Table 76	'H-NMR & (ppm), 300MHz	DMSO-d <sub>6</sub> 1. 58-1. 82(4H, m) 2. 68-2. 84(2H, m) 3. 02-3. 26(2H, m) 3. 67(3H, s) 4. 12-4. 20(2H, m) 4. 82-4. 88(1H, m) 6. 92(1H, d, J=9.01 7. 16-7. 40(8H, m) 7. 54(1H, s) 7. 54(1H, s) 8. 21(1H, s) 8. 21(1H, s) 10. 21(1H, s)	DMSO-d <sub>4</sub> 1. 20(3H, t, J=5. 4 1. 82-1. 85 (4H, m) 2. 58-2. 62(3H, m) 2. 96-3. 04(2H, m) 3. 15-3. 26(2H, m) 4. 11(2H, t, J=4. 3 4. 16(2H, q, J=5. 4 73(2H, d, J=6. 3 7. 73(2H, d, J=6. 3 7. 73(2H, d, J=6. 3 7. 77(2H, d, J=6. 3 8. 72-8. 83(2H, m) 8. 88(1H, d, J=6. 3
30				)Bt
·35			Ph COOMe	CONH COOE
40		Compound	HO COME	
<b>4</b> 5			H <sub>2</sub> N-(CH <sub>2</sub> ), -0	MeN-(CH2),-0- H •HC1
50	•	Bx. No.	146	147

		Ţ	Ţ.
5	Blemental analysis (%)		
10	FAB-MS	467 (free base, MH+)	(free base, MH*)
	[R (cm <sup>-1</sup> )	KBr 3343 2936 1741 1638 1550	KBr 3423 2938 1735 1617 1211
20	m), 300MHz	1=7Hz) 4H. m) 1=6Hz) 1=6Hz) 1=6Hz) 1=6Hz) 1=7Hz) 1=7Hz) 1=7Hz) 1=3. 9Hz) 1=9Hz) 1=9Hz) 1-3.	t. J=6.8Hz) (2(4H. m) (8(6H. m) (2(6H. m) (9, J=6.8Hz) (8(1H. m) (4(5H. m) (4, J=8.6Hz) d, J=8.6Hz) br) d, J=7.2Hz)
Table 77	1H-NMR & (ppm), 300MHz	DMSO-d <sub>6</sub> 1. 16(3H, t, J=' 1. 16(3H, t, J=' 1. 16(3H, t, J=' 1. 16(3H, t, J=' 3. 25(3H, t, J=' 3. 12-3. 25(2H, t, J=' 3. 12-3. 25(2H, t, J=' 7. 54(1H, dd, J=' 7. 59(1H, d, J='	DMSO-d. 1. 14(3H, t. J. 1. 66-1. 82(4) 2. 42-2. 58(6) 2. 84-3. 22(6) 4. 09(2H, q. J. 4. 64-4. 68(11) 7. 18-7. 34(5) 7. 98(2H, d. J. 8. 68(1H, d. J. 9. 04(1H, d. J.
30		- Ph C008t	C00Bt
35	pun	OH CONH	CONH -
40	Compound	N N N	N N N
45		MeN-(CH2), '-	MeN-(CH2), 'HC1
50	Bx. No.	148	149

	Elemental analysis (%)		
10	PAB-MS	439 (free base, MH+)	477 (free base, MH*)
15	IR (cm <sup>-1</sup> )	KBr 1735 1623 1545 1224	KBr 1654 1542 1437 1231
20	pm), 300MHz	38(4H, m) m) 70(6H, m) m) 28(5H, m) 56(2H, m) 6 J=6.0Hz) m) d, J=6.0Hz) H, S)	de H. S. H. M. H. M. H. M. H. M. J=6HZ) H. S. H. d. J=6HZ) H. d. J=6. OHZ) H. d. J=6. OHZ)
75 25 Taple 28.	1H-NMR & (ppm), 300MHz	DMSO-ds 1. 70-1. 88 (4H 2. 53 (3H, m) 2. 90-3. 70 (6H 7. 20-7. 28 (5H 7. 55-7. 56 (2H 8. 04 (1H, d, J= 8. 62 (2H, m) 9. 03 (1H, d, J= 12. 12 (1H, s) 13. 0 (1H, brs)	DMSO-de 1. 69-1. 88 2. 34(3H. s) 2. 33(3H. m) 2. 89-3. 80 5. 62(1H. m) 7. 20-7. 32(7. 52(1H. d. 7. 55(1H. d. 7. 55(1H. d. 7. 55(1H. d. 8. 69(2H. d. 9. 40(1H. d. 12. 06(1H. d.
30		H- H-003	Ph N N Me
35	pun	OH CONH	OH CONH
40	Compound	N N N	N N O
<b>4</b> 5		Men-(CH <sub>2</sub> ), \ H	Men-(CH <sub>2</sub> ), ~ H
50	Š.	150	151

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. 5		Elemental analysis (%)		
15		FAB-MS	442 (free base, MH <sup>+</sup> )	484 (free base, MH*)
20		[R (cm <sup>-1</sup> )		KBr 1738 1643 1497 1469
25 .	Table 79	'H-NMR & (ppm), 300MHz	DMSO-de 1. 15(3H, t, J=15Hz) 2. 18(2H, m) 2. 57(3H, m) 2. 90-3. 50(6H, m) 4. 12(2H, q, J=15Hz) 4. 60(1H, m) 7. 21-7. 34(5H, m) 7. 21-7. 34(5H, m) 7. 82(1H, d, J=9Hz) 8. 99(1H, s) 8. 99(1H, s) 8. 90(2H, s) 8. 91(1H, d, J=6Hz)	DMSO-de 1. 11-1. 83(11H. m) 2. 52(3H. m) 2. 83-3. 57(6H. m) 4. 10(2H, q, J=18Hz) 4. 65(1H. m) 7. 20-7. 83(7H, m) 8. 12(1H. s) 8. 94(3H. m)
35			CONH COOBT	Ph COOB
<b>40</b> .		Compound	N O	N CONH
45			MeN-(CH <sub>2</sub> ) <sub>s</sub> -S- H -HCl	MeN-(CH₂),-S→ H •HC1
50		S	152	153

Table 80

		ignie ou			
RX.	Compound	<sup>1</sup> H-NMR & (ppm), 300MHz	IR (cm <sup>-1</sup> )	IR (cm <sup>-1</sup> ) FAB-MS	Blemental (%)
	OMe Ph	CDCl <sub>3</sub>		445	
	MeN-(CH <sub>8</sub> ),-S-(ONH - C00Bt	1.24(3H, t, J=7.3Hz) 1.70-1.83(2H, m)		MH <sup>+</sup> ).	
		1.97-2.08(2H, m)			
	•HC1	2.65(3H, s) 2.92-3.02(4H, m)			
_		3.21(2H, d, J=5.8Hz)			
154		3.80(3H, s)	, , ,		
}		4.18(2H, q, J=7.3Hz)		٠	
		5.03(1H, q, J=5.8Hz)			
		6.83(1H, d, J=1.3Hz)			
		6.93(1H, dd, J=8.2, 1.3Hz)			
		7.15-7.28(5H, m)			
		8.08(1H, d, J=8.2Hz)			
		8.27(1H, d, J=7.3Hz)			
Ŀ	-	0 560H hm)			

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	8		:	
5	Elemental analysis (			
10	FAB-MS	538 (free base, MH*)	503 (free base, MH+)	
15	IR (cm <sup>-1</sup> )	KBr 3436 1774 1638 1459	KBr 3342 2972 1738 1651 1262 1182	
20	m). 300MHz	=7Hz   4H. m)  =7Hz   =7Hz   H. m)  =9Hz	=6.0Hz   =6.0Hz   H.m   H.m   H.m  =6.0Hz	l. m)
25 q	H-NMR & (ppm), 300MHz	DMSO-d <sub>4</sub> 3. 08-3. 60(1 <sub>4</sub> 4. 12(2H, q, J=7. 15-7. 32(5F) 8. 31(1H, s) 9. 59(2H, dr, J=9. 59(2	0-d (3H, t. J (3H, t. J (3H, t. J (3H, t. J -2. 34(2 -3. 36(6 (2H, q, J	6.98(1H, S) 7.24-7.32(5H, m) 8.11(1H, S) 8.89(1H, brs) 9.08(1H, d, J=7.2
30		Ph COOBt	Ph COOBt	
35	pun	OH CONH	H -CONH	
40	Compound	C1)		
45	·	HN Ŋ-(CH	HP CG	
50	Ex.	155	156	

5 .·	Blemental analysis (%)		
10	FAB-MS B1	(free base, MH <sup>+</sup> )	493 (free base, MH <sup>+</sup> )
	IR (cm <sup>-1</sup> )		ę
20	300MHz	5H2) 5H2)	
rable 82	<sup>1</sup> H-NMR & (ppm), 300MHz	CDC3 2.20-2.34(2H, m) 2.7(3H, s) 2.77(2H, t, J=5Hz) 3.11(2H, brs) 3.23(2H, ddd, J=13, 8, 5Hz) 3.77(3H, s) 5.01(1H, ddd, J=8, 8, 5Hz) 6.60(1H, dd, J=9, 2Hz) 6.73(1H, d, J=9, 2Hz) 7.13(2H, dd, J=8, 2Hz) 7.24-7.32(3H, m) 7.37(1H, d, J=9Hz) 7.37(2H, d, J=9Hz) 7.37(2H, d, J=9Hz) 9.51(2H, s)	CDCl <sub>3</sub> 2.20-2.40(2H, m) 2.71-2.85(4H, m) 3.05-3.25(4H, m) 3.71-1.1. s) 4.9(1H, s) 7.14-7.30(6H, m) 7.67(1H, s) 9.43(2H, brs) 11.91(1H, s)
30 E	N-H <sub>I</sub>	CDC <sub>13</sub> 2.20-2.34(2H, m) 2.71(3H, s) 2.77(2H, t, J=5Hz) 3.11(2H, brs) 3.23(2H, ddd, J=1:3.77(3H, s) 5.01(1H, ddd, J=9, 6.60(1H, dd, J=9, 6.73(1H, d, J=8Hz) 7.13(2H, d, J=8Hz) 7.13(2H, d, J=8Hz) 7.13(2H, d, J=9Hz) 7.24-7.32(3H, m) 7.34(1H, d, J=9Hz) 7.37(1H, d, J=9Hz)	CDCl <sub>3</sub> 2.20-240(2H, m) 2.71-2.85(4H, m) 3.05-3.25(4H, m) 3.05-3.25(4H, m) 3.71(1H, s) 4.9(1H, q, J=7.2H, 6.86(1H, s) 7.14-7.30(6H, m) 7.67(1H, s) 9.43(2H, brs) 11.91(1H, s)
35		- COOMe	Ph COOMe
40	Compound	OTH CONF	Br OH
45	·	Men-(CH <sub>8</sub> ) <sub>8</sub> -C00 H -HC1	MeN-(CH1),-C00 — H Br
50 .	RX. No.		158 Me

Formulation Examples of the pharmaceutical agents containing the compound of the present invention are shown in th following.

## Formulation Example 1 (Tablet)

	4				
ı	:	1	ı		

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	_
(1) Compound of Example 18	10 g
(2) Lactose	50 g
(3) Corn starch	15 g
(4) Carboxymethylcellulose sodium	44 g
(5) Magnesium stearate	1 g

The entire amounts of (1), (2) and (3), and 30 g of (4) were kneaded with water, dried in vacuo, and granulated. The granules were added with 14 g of (4) and 1 g of (5), and the mixture was compressed to give tablets, whereby 1,000 tablets containing 10 mg of the compound per tablet were prepared.

### Formulation Example 2 (Injection)

The compound of Example 18 (100 mg) was dissolved in an aqueous solution of mannitol (5 g) dissolved in water (100 ml) for injection, sterilized by filtration through a 0.22 µm filter, and filled in sterilized ampoules by 1 ml to give injections containing 1 mg of the compound per ampoule.

The results of experiments with respect to the suppression of production of inflammatory cytokines, suppression of LPS-induced peritonitis and suppression of LPS/D-glactosamine-induced hepatitis by the compound of the present invention are shown below.

### Experimental Example 1: Suppression of production of inflammatory cytokines

Thirty ml of human peripheral blood added with heparin was placed on Ficol-Paque (15 ml), and centrifuged at 400 G for 40 minutes at room temperature. The obtained monocyte fraction layers were collected and washed three times with E-MEM medium. The cells were adjusted to a final concentration of  $0.5 \times 10^5$  cells/800  $\mu$ l with RPMI-1640 medium containing 5% bovine fetal serum (2-mercaptoethanol), and seeded in a 24 well plate by 800  $\mu$ l. A test sample (100  $\mu$ l) was added and 100  $\mu$ l of lipopolysaccharide (LPS, 100  $\mu$ g/ml) was added one hour later. The supernatant was taken at 20 hours after stimulation with LPS, and amounts of various cytokines were determined using an ELISA kit. By plotting the cytokine amounts at various concentrations, the concentration of the test sample necessary for inhibition by 50% (IC<sub>50</sub>) was determined. The results are shown in Tables 83-88.

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#### Table 83

IC <sub>50</sub> (μM)		
IL-1β	TNF	IL-8
0.002	0.008	0.009
-	-	0.01
>30	14	>30
3	2	2
75	6	6
14	6	14
•	•	8
-	•	<0.3
•	-	0.6
-	-	0.4
•	-	1
	0.002 - >30 3 75	IL-1β TNF 0.002 0.008  >30 14 3 2 75 6

## Table 83 (continued)

IC<sub>50</sub> (μM) TNF IL-1β IL-8 1 Example No. 15 0.03 Example No. 16 <0.01 Example No. 18 <0.01 Example No. 19 29 Example No. 20 Example No. 21 <0.01 Example No. 22 <0.01 0.02 Example No. 24 -Example No. 25 0.01 Example No. 26 0.009

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Table 84

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	IC <sub>50</sub> (μM)		
	IL-1β	TNF	IL-8
Example No. 27		-	<0.01
Example No. 28	-	•	<0.01
Example No. 29	-	-	<0.01
Example No. 30	-		0.6
Example No. 31	-	-	<0.01
Example No. 32	-	•	0.5
Example No. 34	-	•	2
Example No. 36	-	•	0.06
Example No. 37	•	•	0.3
Example No. 39	-	•	0.02
Example No. 40	-	-	0.01
Example No. 41	•	•	<0.01
Example No. 42	-	-	0.1
Example No. 43	•	•	0.03
Example No. 44	-	·	<0.01
Example No. 45	0.0008	0.004	0.004
Example No. 46	-		<0.01
Example No. 47	•	•	3
Example No. 48		-	0.2
Example No. 49	_		0.02
Example No. 50	_	<u> </u>	28

Table 85

	IC <sub>50</sub> (μM)		
	IL-1β	TNF	IL-8
Example No. 51	-	-	7
Example No. 52	-	-	<0.01
Example No. 53	-	-	<0.01
Example No. 54	-	-	<0.01
Example No. 55	-	-	<0.01
Example No. 56	-	-	4
Example No. 57	-		0.05
Example No. 58	-	-	0.02
Example No. 60	-	-	0.03
Example No. 63	-	•	0.1
Example No. 64	-	-	0.05
Example No. 67	-	-	0.05
Example No. 68	-	-	0.001
Example No. 69	-	-	<0.001
Example No. 70	-	-	0.006
Example No. 71	•	-	0.04
Example No. 72	-	-	0.1
Example No. 73	-	-	<0.01
Example No. 74		-	0.07
Example No. 75	•	-	0.04
Example No. 76	-	-	0.3

Table 86

	IC <sub>50</sub> (μM)		
	IL-1β	TNF	IL-8
Example No. 77		-	3
Example No. 80	-	-	3
Example No. 81	-	-	4
Example No. 82	-	-	0.02
Example No. 83	-	-	0.09
Example No. 84	-	-	0.03
Example No. 85	-	-	0.07
Example No. 86	-	-	<0.001

## Table 86 (continued)

IC<sub>50</sub> (μM) IL-1β TNF IL-8 Example No. 87 0.2 Example No. 88 3 Example No. 89 -0.6 Example No. 90 0.6 Example No. 91 0.001 Example No. 92 0.03 Example No. 94 1 Example No. 95 0.09 Example No. 96 0.003 Example No 98 0.001 Example No. 99 0.001 Example No. 100 0.001 --Example No. 101 0.003

Table 87

	IC <sub>50</sub> (μM)		
	IL-1β	TNF	IL-8
Example No. 102	-	-	0.002
Example No. 103	-	•	0.7
Example No. 104	-	•	0.7
Example No. 105	0.001	0.004	0.005
Example No. 106	-	-	<0.01
Example No. 110	-	-	<0.01
Example No. 111	-		<0.01
Example No. 117	-	•	<0.01
Example No. 122	-	•	<0.01
Example No. 125	-	•	0.01
Example No. 126	-	•	0.8
Example No. 127	-	•	0.2
Example No. 128	-	•	0.2
Example No. 129	-	•	2
Example No. 132	•	-	0.07
Example No. 133	-	-	0.2
Example No. 134	•	-	0.2
Example No. 136	-	-	0.2
Example No. 137	•	-	2

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Table 87 (continued)

	IC <sub>50</sub> (μM)			
	IL-1β TNF IL-8			
Example No. 138		-	1	
Example No. 139	•	•	4	

Table 88

	IC <sub>50</sub> (μM)		
	IL-1β	TNF	IL-8
Example No. 140	-	-	13
Example No. 141	-	-	3
Example No. 142	-	-	0.4
Example No. 143	-	-	3
Example No. 144	-	-	29
Example No. 146	-	-	5
Example No. 147	-	-	2
Example No. 148	-	-	4
Example No. 149	٠	-	3
Example No. 152	•	-	7
Example No. 153	-	-	1
Example No. 155	-	-	0.2
Example No. 156	-	-	2

Experimental Example 2: Suppression of LPS-induced peritonitis

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LPS (30  $\mu$ g/ml, 1 ml) prepared with physiological saline containing 0.5% CMC (carboxymethylcellulose) was intraperitoneally injected into male Balb/c mice to induce peritonitis. One hour later, the mice were killed with carbon dioxide, and the amount of TNF  $\alpha$  in the peritoneal fluid was determined using an ELISA kit.

The test sample (50 mg/kg) was administered from the tail vein at 60 minutes before LPS injection, and the degree of suppression was investigated. The suppression by the test sample is shown in the ratio relative to the suppression in the control group.

Suppression (%) =  $100 \cdot \text{(TNF amount of group treated with test sample/TNF amount of control group)} \times 100$ 

The results are shown in Table 89 wherein \*\* means the presence of significant difference by p<0.01 from the control group.

Table 89

	Inhibition (%)
Example No. 1	64**
Example No. 4	38**
Example N . 9	21

Table 89 (continued)

	Inhibition (%)
Example No. 19	32**
Example No. 51	38**
Example No. 52	31**
Example No. 148	19
Example No. 155	28**

Experimental Example 3: Suppression of LPS/D-galactosamine-induced hepatitis

LPS (5 μg/kg)/D-galactosamine (500 mg/kg) in physiological saline was intraperitoneally injected to male C57BL/6 mice to induce hepatitis. Six hours after the injection of LPS/D-galactosamine in physiological saline, blood was taken from the mice orbital venosus plexus. Plasma was separated from the blood, and ALT in blood was determined by a biochemical analyzer. The test sample was administered from the tail vein at 10 minutes before the injection of LPS/D-galactosamine in physiological saline, and the degree of suppression was investigated. The suppression by the test sample is shown in the ratio relative to the suppression in the control group.

Suppression (%) =  $100 - (ALT amount of group treated with test sample/ALT amount of control group) <math>\times 100$ 

The results are shown in Tables 90-91.

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Table 90

	Dose (mg/kg)	Inhibition (%)
Example No. 1	5	88
	10	78
Example No. 2	10	65
Example No. 4	10	42
Example No. 10	10	77
Example No. 18	10	86
Example No. 22	10	. 51
Example No. 24	10	63
Example No. 27	10	67
Example No. 31	5	87
Example No. 32	10	78
Example No. 36	10	47
Example No. 37	10	80

Table 91

	Dose (mg/kg)	Inhibition (%)
Example No. 40	10	49
Example No. 45	10	30
Example No. 46	10	57

Table 91 (continued)

	Dose (mg/kg)	Inhibition (%)
Example No. 50	10	15
Example No. 54	5	74
Example No. 60	10	42
Example No. 61	10	6
Example No. 105	10	40
Example No. 117	10	54
Example No. 123	10	41
Example No. 126	10	27
Example No. 127	5	82
Example No. 138	5	22

From the foregoing results, it is evident that the compound of the present invention suppresses production of inflammatory cytokines and is useful for the prophylaxis and therapy of noninfectious or infectious diseases accompanied by neutrophile migration, which are represented by rheumatic diseases (e.g., rheumatoid arthritis); arthritis due to gout; systemic lupus erythematosus; dermatopathy (e.g., psoriasis, pustulosis and atopic dermatitis); respiratory diseases (e.g., bronchial asthma, bronchitis, ARDS and diffused interstitial pneumonia); inflammatory bowel diseases (e.g., ulcerative colitis and Crohn's disease); acute or chronic hepatitis inclusive of fulminant hepatitis; acute or chronic glomerulonephritis; nephropyelitis; uveitis caused by Behcet disease and vogt-Koyanagi Harada disease; Mediterranean fever (polyserositis); ischemic diseases (e.g., myocardial infarction); and systemic circulatory failure and multiorgan failure caused by sepsis.

The test results with respect to inflammatory cytokines such as IL-6 and GM-CSF have confirmed suppression of these inflammatory cytokines by the compound of the present invention.

### Claims

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### 1. An amide compound of the formula (I):

$$R - A - X \xrightarrow{R^1 \qquad R^2 \qquad 0} \qquad (CH_2)_m \qquad R^6$$

$$R - A - X \xrightarrow{R^3 \qquad R^4 \qquad R^5} \qquad (I)$$

wherein;

R

Α

X

is an optionally substituted non-aromatic heterocyclic group containing nitrogen, a hydroxy,  $R_{\rm a}$ , an alkoxy substituted by  $R_{\rm a}$ , an alkylthio substituted by  $R_{\rm a}$ , or an alkylamino substituted by  $R_{\rm b}$ 

wherein  $R_a$  is amino, guanidino, amidino, carbamoyl, ureido, thioureido, hydrazino, hydrazino-carbonyl or imino, these groups being optionally substituted by a substituent selected from the group consisting of lower alkyl, halogenated lower alkyl, cycloalkyl, aralkyl, aryl and amino-protecting group;

is an optionally substituted, linear or branched alkylene which optionally has one or more double bond(s) or triple bond(s) in the chain, or a single bond:

is an oxygen atom, a sulfur atom, a cycloalkylene, a divalent aromatic heterocyclic group having one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, -SO-, -SO<sub>2</sub>-, -C=C-, -C=C-, -CO-, -COO-, -OOC-, -CS-, -COS-, -O-CO-O-, -NH-CO-NH-, -NH-CS-NH-, -NH-C(=NH)-NH-, -NR<sup>8</sup>-, -NR<sup>8</sup>CO-, -CONR<sup>8</sup>-, -NR<sup>8</sup>SO<sub>2</sub>-, -SO<sub>2</sub>NR<sup>8</sup>-, -NR<sup>8</sup>-COO-, -OOC-NR<sup>8</sup>-, or -CR<sup>9</sup>R<sup>10</sup>-

		wherein R <sup>8</sup> is hydrogen atom, alkyl, cycloalkyl, aryl, aralkyl or amino-protecting group, and R <sup>9</sup> and R <sup>10</sup> are the same or different and each is hydrogen atom, alkyl, cycloalkyl, aryl or aralkyl;
	M	is an arylene, a cycloalkylene, or a divalent heterocyclic group which has one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom,
5		and which optionally forms a fused ring;
	$R^1$ , $R^2$ , $R^3$ and $R^4$	are the same or different and each is a hydrogen atom, a hydroxy, a halogen atom, an alkoxy,
		a mercapto, an alkylthio, a nitro, a cyano, a carboxy, an alkoxycarbonyl, an aryloxycarbonyl,
		an acyl, an alkyl optionally substituted by a substituent selected from the group consisting of
		hydroxy, lower alkoxy and halogen atom, an amino optionally substituted by a substituent
10		selected from the group consisting of alkyl, aryl, aralkyl and amino-protecting group, or -O-CO-R <sup>11</sup>
		wherein R <sup>11</sup> is optionally substituted alkoxy, optionally substituted aryl, optionally substituted
		cycloalkyl, optionally substituted aryloxy, optionally substituted aralkyloxy, optionally substi-
		tuted alkylthio, optionally substituted arylthio, or alkyl optionally substituted by a substituent selected from the group consisting of alkoxycarbonyl, acyloxy, aryl, aryloxy, aryloxycarbonyl,
15		aralkyloxy, aralkyloxycarbonyl, alkylthio, arylthio, acyl, lower alkoxy, carboxy, halogen atom
		and amino optionally substituted by lower alkyl or acyl;
	R <sup>5</sup>	is a hydrogen atom, an alkyl optionally substituted by a halogen atom, an optionally substi-
	••	tuted aralkyl, or an amino-protecting group;
20	m	is 0 or an integer of 1-6;
	R <sup>6</sup>	is an optionally substituted aryl, an optionally substituted cycloalkyl, an optionally substituted
		lower alkyl, an optionally substituted lower alkoxy, an optionally substituted lower alkylthio, an
		amino optionally substituted by a substituent selected from the group consisting of lower alkyl,
		aryl, aralkyl and amino-protecting group, or an optionally substituted heterocyclic group hav-
25		ing one ore more hetero atoms selected from the group consisting of a nitrogen atom, sulfur
	R <sup>7</sup>	atom and oxygen atom; and is a hydrogen atom, an optionally substituted alkyl, an optionally substituted aryl, an optionally
	n	substituted aromatic heterocyclic group having one ore more hetero atoms selected from the
		group consisting of a nitrogen atom, sulfur atom and oxygen atom,
30		or -CO(Y) <sub>0</sub> R <sup>12</sup>
		wherein Y is oxygen atom, sulfur atom, -NR <sup>13</sup> - or -NR <sup>13</sup> -SO <sub>2</sub> -wherein R <sup>13</sup> is hydrogen atom,
		alkyl, aralkyl, hydroxy, alkoxy, aryl or amino-protecting group, p is 0 or 1, and R <sup>12</sup> is hydrogen
		atom, optionally substituted alkenyl, optionally substituted alkynyl, optionally substituted
		cycloalkyl, optionally substituted aryl, optionally substituted aralkyl, adamantyl, cycloalkylide-
35		neamino, optionally substituted heterocyclic group having one or more hetero atom(s) selected from the group consisting of nitrogen atom, sulfur atom and oxygen atom, or alkyl
		optionally substituted by a substituent selected from the group consisting of hydroxy, alkoxy,
		alkoxyalkoxy, alkoxycarbonyl, acyloxy, carboxy, heterocyclic group having one or more hetero
		atom(s) selected from the group consisting of nitrogen atom, sulfur atom and oxygen atom,
40		and amino optionally substituted by a substituent selected from the group consisting of alkyl,
		aryl, aralkyl and amino-protecting group;
	or a pharmass for	ally acceptable acid addition salt thereof.
	or a priarmaceutica	any acceptable add addition sait thereof.

2. The amide compound of claim 1, wherein, in the formula (I), at least one symbol selected from the group consisting of R, A, X, M, R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup>, R<sup>4</sup>, R<sup>5</sup>, m, R<sup>6</sup> and R<sup>7</sup> satisfies the following definitions, or a pharmaceutically acceptable acid addition salt thereof:

	R	is a non-aromatic heterocyclic group containing nitrogen, which is optionally substituted by
50		lower alkyl or amino-protecting group, Ra1, an alkoxy substituted by Ra1, an alkylthio substi-
		tuted by R <sub>a1</sub> , or an alkylamino substituted by R <sub>a1</sub> ,
		wherein Rat is amino, guanidino, amidino, carbamoyl, ureido, thioureido, hydrazino, hydrazi-
		nocarbonyl or imino, these groups being optionally substituted by a substituent selected from
		the group consisting of lower alkyl, aralkyl and amino-protecting group;
55	Α	is a linear or branched alkylene which optionally has one or more double bond(s) or triple
		bond(s) in the chain, or a single bond;
	X	is an oxygen atom, a sulfur atom, a cycloalkylene, a divalent aromatic heterocyclic group hav-
		ing one or mor hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur

atom and oxygen atom, -SO-, -SO<sub>2</sub>-, -C=C-, -C=C-, -CO-, -COO-, -OC-, -CS-, -COS-, -O-CO-O-, -NH-CO-NH-, -NH-CS-NH-, -NH-C(=NH)-NH-, -NR8-, -NR8-CO-, -CONR8-, NR8-SO2-, -SO2NR8-, -NR8-COO-, -OOC-NR8-, or -CR9-R10wherein R8' is hydrogen atom, lower alkyl, aralkyl or amino-protecting group, and R9' and R10' are the same or different and each is hydrogen atom, lower alkyl or aralkyl; 5 М is an arylene, a cycloalkylene or a divalent heterocyclic group which has one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, and which optionally forms a fused ring; R1, R2, R3 and R4 are the same or different and each is a hydrogen atom, a hydroxy, a halogen atom, a lower 10 alkoxy, a mercapto, a lower alkytthio, a nitro, a cyano, a carboxy, a lower alkoxycarbonyl, an aryloxycarbonyl, an acyl, a lower alkyl optionally substituted by a substituent selected from the group consisting of hydroxy, lower alkoxy and halogen atom, an amino optionally substituted by a substituent selected from the group consisting of lower alkyl, aralkyl and amino-protecting group, or -O-CO-R111 15 wherein R111 is lower alkoxy, optionally substituted cycloalkyl, lower alkyl optionally substituted tuted by a substituent selected from the group consisting of lower alkoxycarbonyl, acyloxy, aralkyloxy, aralkyloxycarbonyl, acyl, lower alkoxy, carboxy and amino optionally substituted by lower alkyl, or aryl optionally substituted by a substituent selected from the group consisting 20 of lower alkyl, carboxy and benzyloxycarbonyl; R<sup>5</sup> is a hydrogen atom, an alkyl optionally substituted by a halogen atom, an optionally substituted aralkyl, or an amino-protecting group: is 0 or an integer of 1-6; R<sup>6</sup> is an aryl, a cycloalkyl, or a heterocyclic group having one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom 25 wherein said aryl, cycloalkyl and heterocyclic group having one or more hetero atom(s) selected from the group consisting of nitrogen atom, sulfur atom and oxygen atom are optionally substituted by a substituent selected from the group consisting of lower alkyl, halogen atom, hydroxy, lower alkoxy, amino, carboxy and lower alkoxycarbonyl; and R<sup>7</sup> is a hydrogen atom, a lower alkyl optionally substituted by a substituent selected from the 30 group consisting of hydroxy, lower alkoxy, mercapto, lower alkylthio, carboxy, lower alkoxycarbonyl and amino, an aromatic heterocyclic group which has one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, and which is optionally substituted by lower alkyl, or -CO (Y)<sub>n</sub>R<sup>12</sup> wherein Y is oxygen atom, sulfur atom, -NR13'- or -NR13'-SO2-35 wherein R131 is hydrogen atom, lower alkyl, aralkyl, hydroxy, lower alkoxy or amino-protecting p is 0 or 1, and R12, is hydrogen atom, aralkyl, adamantyl, cycloalkylideneamino, cycloalkyl optionally substituted by lower alkyl, alkyl optionally substituted by a substituent selected from 40 the group consisting of hydroxy, lower alkoxy, lower alkoxy lower alkoxy, lower alkoxycarbonyl, acyloxy, carboxy, heterocyclic group having one or more hetero atom(s) selected from the group consisting of nitrogen atom, sulfur atom and oxygen atom, and amino optionally substituted by a substituent selected from the group consisting of lower alkyl, aralkyl and amino-protecting group, aryl optionally substituted by a substituent selected from the group consisting 45 of lower alkyl, halogen atom, amino, carboxy, hydroxy and lower alkoxy, or heterocyclic group which is optionally substituted by a substituent selected from the group consisting of lower alkyl, halogen atom, amino, carboxy, hydroxy and lower alkoxy, and which has one or more hetero atom(s) selected from the group consisting of nitrogen atom, sulfur atom and oxygen atom. 50 The amide compound of claim 1, wherein, in the formula (I), at least one symbol selected from the group consisting of R, A, X, M, R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup>, R<sup>4</sup>, R<sup>5</sup>, m, R<sup>6</sup> and R<sup>7</sup> satisfies the following definitions, or a pharmaceutically acceptable

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tuted by lower alkyl or amino-protecting group:

lower alkyl or amino-protecting group, R<sub>a2</sub>, or an alkoxy substituted by R<sub>a2</sub>,

is a non-aromatic heterocyclic group containing nitrogen, which is optionally substituted by

wherein R<sub>a2</sub> is amino, guanidino, amidino or carbamoyl, these groups being optionally substi-

acid addition salt thereof:

R

	٨	is a linear alkylene or a single bond;
	A	
	X	is an oxygen atom, a sulfur atom, a divalent aromatic heterocyclic group having one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, -COO-, -OOC-, -NR8"-, -NR8"CO-, -CONR8"-, -NR8"SO <sub>2</sub> -, -SO <sub>2</sub> NR8"-, or -CR9"R10"-
5		wherein R <sup>8</sup> " is hydrogen atom, lower alkyl or amino-protecting group, and R <sup>9</sup> " and R <sup>10</sup> " are the same or different and each is hydrogen atom or lower alkyl;
	М	is an arylene or a divalent heterocyclic group which has one or more hetero atom(s) selected
	141	from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, and which option- ally forms a fused ring;
10	$R^1$ , $R^2$ , $R^3$ and $R$	are the same or different and each is a hydrogen atom, a hydroxy, a halogen atom, lower alkoxy, a lower alkyl optionally substituted by a substituent selected from the group consisting of hydroxy, lower alkoxy and halogen atom, or -O-CO-R <sup>11</sup> "
		wherein R <sup>11</sup> " is lower alkoxy, cycloalkyl, aryl optionally substituted by lower alkyl, or lower alkyl optionally substituted by a substituent selected from the group consisting of acyloxy,
45		aralkyloxycarbonyl and amino optionally substituted by lower alkyl;
15	R <sup>5</sup>	is a hydrogen atom, a lower alkyl, or an amino-protecting group;
		is 1;
	m R <sup>6</sup>	is an aryl or a cycloalkyl
	n	wherein said aryl and cycloalkyl are optionally substituted by halogen atom or hydroxy; and
20	R <sup>7</sup>	is a hydrogen atom, a lower alkyl optionally substituted by hydroxy or lower alkoxy, an aromatic heterocyclic group which has one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, and which is optionally substituted by lower alkyl, or
		-CO(Y") <sub>p</sub> R <sup>12</sup> "
25		wherein Y" is oxygen atom, sulfur atom or -NR <sup>13</sup> "-
		wherein R <sup>13</sup> " is hydrogen atom, lower alkyl, hydroxy or amino-protecting group, p is 0 or 1,
		and R12" is hydrogen atom, aralkyl, adamantyl, cycloalkylideneamino, cycloalkyl optionally
		substituted by lower alkyl, aryl optionally substituted by halogen atom, alkyl optionally substi-
		tuted by a substituent selected from the group consisting of hydroxy, lower alkoxy, lower alkoxy
30		lower alkoxy, lower alkoxycarbonyl, acyloxy, carboxy, heterocyclic group having one or more hetero atom(s) selected from the group consisting of nitrogen atom, sulfur atom and oxygen atom, and amino optionally substituted by a substituent selected from the group consisting of
		lower alkyl, aralkyl and amino-protecting group, or heterocyclic group which is optionally sub-
		stituted by lower alkyl, and which has one or more hetero atom(s) selected from the group
35		consisting of nitrogen atom, sulfur atom and oxygen atom.
	4. The amide comp of R, A, X, M, R <sup>1</sup> acid addition sal	ound of claim 1, wherein, in the formula (I), at least one symbol selected from the group consisting , R <sup>2</sup> , R <sup>3</sup> , R <sup>4</sup> , R <sup>5</sup> , m, R <sup>6</sup> and R <sup>7</sup> satisfies the following definitions, or a pharmaceutically acceptable
40	acid addition sai	Thereof.
40	R	is a piperazinyl optionally substituted by lower alkyl, a piperidyl optionally substituted by lower alkyl, an amino, or a lower alkoxy substituted by amino wherein amino is optionally substituted by lower alkyl;
	٨	is a linear alkylene;
45	A X	is an oxygen atom, a sulfur atom, -NH- or -CH <sub>2</sub> -;
45	M	is an arylene;
	R <sup>1</sup> , R <sup>2</sup> , R <sup>3</sup> and I	R <sup>4</sup> are the same or different and each is a hydrogen atom, a hydroxy, a halogen atom, or -O-CO-R <sup>11</sup>
		wherein R11m is lower alkyl optionally substituted by a substituent selected from the group
50		consisting of amino, acyloxy and benzyloxycarbonyl, or phenyl optionally substituted by lower alkyl;
	R <sup>5</sup>	is a hydrogen atom;
	m	is 1;
	R <sup>6</sup>	is a phenyl; and
55	R <sup>7</sup>	is -COO-R <sup>12</sup> '''
		wherein R <sup>12</sup> " is hydrogen atom, aralkyl, adamantyl, cyclohexylideneamino, cyclohexyl optionally substituted by lower alkyl, piperidyl optionally substituted by lower alkyl, or alkyl optionally substituted by a substituent selected from the group consisting of hydroxy, lower alkoxy, lower
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alkoxy lower alkoxy, lower alkoxycarbonyl, acyloxy, piperazinyl and amino optionally substituted by lower alkyl.

- The amide compound of claim 4, wherein M is phenylene, or a pharmaceutically acceptable acid addition salt thereof.
- The amide compound of claim 4, wherein R<sup>7</sup> is -COO-R<sup>12-m</sup> wherein R<sup>12-m</sup> is lower alkyl, or cyclohexyl which is
  optionally substituted by lower alkyl, or a pharmaceutically acceptable acid addition salt thereof.
- 7. The amide compound of claim 4, wherein X is an oxygen atom or -CH<sub>2</sub>-, or a pharmaceutically acceptable acid addition salt thereof.
  - The amide compound of claim 4, wherein R<sup>6</sup> is phenyl and m is 1, or a pharmaceutically acceptable acid addition salt thereof.
  - The amide compound of claim 4, wherein R is amino optionally substituted by lower alkyl, piperazinyl optionally substituted by lower alkyl, or piperidyl optionally substituted by lower alkyl, or a pharmaceutically acceptable acid addition salt thereof.
- 10. The amide compound of claim 4, wherein R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup> and R<sup>4</sup> are the same or different and each is a hydrogen atom, hydroxy, a halogen atom, or -O-CO-R<sup>11</sup>" wherein R<sup>11</sup>" is lower alkyl or phenyl, or a pharmaceutically acceptable acid addition salt thereof.
  - 11. A carboxylic acid compound of the formula (I-a)

$$R \longrightarrow A \longrightarrow X \longrightarrow M \longrightarrow COOH$$
 (I-a)

wherein;

X

M

R1, R2, R3 and R4

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35 R is an optionally substituted non-aromatic heterocyclic group containing nitrogen, a hydroxy, R<sub>a</sub>, an alkoxy substituted by R<sub>a</sub>, an alkylthio substituted by R<sub>a</sub>, or an alkylamino substituted by R<sub>a</sub>,

wherein  $R_a$  is amino, guanidino, amidino, carbamoyl, ureido, thioureido, hydrazino, hydrazino-carbonyl or imino, these groups being optionally substituted by a substituent selected from the group consisting of lower alkyl, halogenated lower alkyl, cycloalkyl, aralkyl, aryl and amino-protecting group;

A is an optionally sutstituted, linear or branched alkylene which optionally has one or more double bond(s) or triple bond(s) in the chain, or a single bond;

is an oxygen atom, a sulfur atom, a cycloalkylene, a divalent aromatic heterocyclic group having one ore more hetero atoms selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, -SO-, -SO<sub>2</sub>-, -C=C-, -C=C-, -CO-, -COO-, -OOC-, -CS-, -COS-, -O-CO-O-, -NH-CO-NH-, -NH-CS-NH-, -NH-C(=NH)-NH-, -NR<sup>8</sup>-, -NR<sup>8</sup>CO-, -CONR<sup>8</sup>-, -NR<sup>8</sup>SO<sub>2</sub>-, -SO<sub>2</sub>NR<sup>8</sup>-, -NR<sup>8</sup>-COO-, -OOC-NR<sup>8</sup>-, or -CR<sup>9</sup>R<sup>10</sup>-

wherein R<sup>8</sup> is hydrogen atom, alkyl, cycloalkyl, aryl, aralkyl or amino-protecting group, and R<sup>9</sup> and R<sup>10</sup> are the same or different and each is hydrogen atom, alkyl, cycloalkyl, aryl or aralkyl; is an arylene, a cycloalkylene or a divalent heterocyclic group which has one ore more hetero atoms selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, and which optionally forms a fused ring; and

are the same or different and each is a hydrogen atom, a hydroxy, a halogen atom, an alkoxy, a mercapto, an alkylthio, a nitro, a cyano, a carboxy, an alkoxycarbonyl, an acyl, an alkyl optionally substituted by a substituent selected from the group consisting of hydroxy, lower alkoxy and halogen atom, an amino optionally substituted by a substituent selected from the group consisting of alkyl, aryl, aralkyl and amino-protecting group, or -O-

CO-R11

wherein R<sup>11</sup> is optionally substituted alkoxy, optionally substituted aryl, optionally substituted cycloalkyl, optionally substituted aryloxy, optionally substituted aralkyloxy, optionally substituted aryloxy, optionally substituted aryloxy, optionally substituted by a substituted substituted by a substituted substituted by a substituted selected from the group consisting of alkoxycarbonyl, acyloxy, aryloxy, aryloxy, aryloxycarbonyl, aralkyloxy, aralkyloxycarbonyl, alkylthio, arylthio, acyl, lower alkoxy, carboxy, halogen atom and amino optionally substituted by lower alkyl or acyl.

12. The carboxylic acid compound of claim 11, wherein, in the formula (I-a), at least one of R, A, X, M, R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup> and R<sup>4</sup> satisfies the following definitions:

is a piperazinyl optionally substituted by lower alkyl, a piperidyl optionally substituted by lower

alkyl, an amino or a lower alkoxy substituted by amino wherein amino is optionally substituted

by lower alkyl;

15 A is a linear alkylene;

X is an oxygen atom, a sulfur atom, -NH- or CH<sub>2</sub>-;

M is an arylene; and R1, R2, R3 and R4 are the same or different and each is a hydrogen atom, a hydroxy, a halogen atom, or -O-CO-

R11"

wherein R<sup>11</sup>" is a lower alkyl optionally substituted by a substituent selected from the group consisting of amino, acyloxy and benzyloxycarbonyl, or phenyl optionally substituted by lower alkyl.

13. An amide compound of the formula (I-b)

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R

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M

is an oxygen atom, a sulfur atom, a cycloalkylene, a divalent aromatic heterocyclic group having one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, -SO-, -SO $_2$ -, -C=C-, -C=C-, -CO-, -COO-, -OOC-, -CS-, -COS-, -O-CO-O-, -NH-CO-NH-, -NH-CS-NH-, -NH-C(=NH)-NH-, -NR $^8$ -, -NR $^8$ CO-, -CONR $^8$ -, -NR $^8$ SO $_2$ -, -SO $_2$ NR $^8$ -, -NR $^8$ -COO-, -OOC-NR $^8$ - or -CR $^9$ R $^{10}$ -

wherein R<sup>8</sup> is hydrogen atom, alkyl, cycloalkyl, aryl, aralkyl or amino-protecting group, and R<sup>9</sup> and R<sup>10</sup> are the same or different and each is hydrogen atom, alkyl, cycloalkyl, aryl or aralkyl; is an arylene, cycloalkylene, or a divalent heterocyclic group which has one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, and which optionally forms a fused ring;

R1, R2, R3 and R4

are the same or different and each is a hydrogen atom, a hydroxy, a halogen atom, an alkoxy, a mercapto, an alkylthio, a nitro, a cyano, a carboxy, an alkoxycarbonyl, an aryloxycarbonyl, an acyl, an alkyl optionally substituted by a substituent selected from the group consisting of hydroxy, lower alkoxy and halogen atom, an amino optionally substituted by a substituent selected from the group consisting of alkyl, aryl, aralkyl and amino-protecting group, or -O-CO-R<sup>11</sup>

CO-F

wherein R<sup>11</sup> is optionally substituted alkoxy, optionally substituted aryl, optionally substituted cycloalkyl, optionally substituted aryloxy, optionally substituted aralkyloxy, optionally substituted alkylthio, optionally substituted arylthio, or alkyl optionally substituted by a substitutent selected from the group consisting of alkoxycarbonyl, acyloxy, aryl, aryloxy, aryloxycarbonyl, aralkyloxy, aralkyloxycarbonyl, alkylthio, arylthio, acyl, lower alkoxy, carboxy, halogen atom and amino optionally substituted by lower alkyl or acyl;

 $R^5$ 

is a hydrogen atom, an alkyl optionally substituted by a halogen atom, optionally substituted

aralkyl, or an amino-protecting group;

is 0 or an integer of 1-6;  $R^6$ 

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R<sup>7</sup>

is an optionally substituted aryl, an optionally substituted cycloalkyl, an optionally substituted lower alkyl, an optionally substituted lower alkoxy, an optionally substituted lower alkylthio, an amino optionally substituted by a substituent selected from the group consisting of lower alkyl, aryl, aralkyl and amino-protecting group, or an optionally substituted heterocyclic group having one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur

atom and oxygen atom; and

is a hydrogen atom, an optionally substituted alkyl, an optionally substituted aryl, an optionally substituted aromatic heterocyclic group having one or more hetero atom(s) selected from the

group consisting of a nitrogen atom, sulfur atom and oxygen atom, or -CO(Y), R12

wherein Y is oxygen atom, sulfur atom, -NR13- or -NR13-SO2-wherein R13 is hydrogen atom, alkyl, aralkyl, hydroxy, alkoxy, aryl or amino-protecting group, p is 0 or 1, and R12 is hydrogen atom, optionally substituted alkenyl, optionally substituted alkynyl, optionally substituted cycloalkyl, optionally substituted aryl, optionally substituted aralkyl, adamantyl, cycloalkylideneamino, alkyl optionally substituted by a substituent selected from the group consisting of hydroxy, alkoxy, alkoxyalkoxy, alkoxycarbonyl, acyloxy, carboxy, heterocyclic group having one or more hetero atom(s) selected from the group consisting of nitrogen atom, sulfur atom and oxygen atom, and amino optionally substituted by a substituent selected from the group consisting of alkyl, aryl, aralkyl and amino-protecting group, or optionally substituted heterocyclic group having one or more hetero atom(s) selected from the group consisting of nitrogen atom, sulfur atom and oxygen atom.

14. The amide compound of claim 13, wherein, in the formula (I-b), at least one symbol selected from the group consisting of X, M, R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup>,

R4, R5, m, R6 and R7 satisfies the following definitions:

is an oxygen atom, a sulfur atom or -NH-;

M is an arylene:

R1, R2, R3 and R4 are the same or different and each is a hydrogen atom, a hydroxy, a halogen atom, or -O-

wherein R11m is lower alkyl optionally substituted by a substituent selected from the group consisting of amino, acyloxy and benzyloxycarbonyl, or a phenyl optionally substituted by

lower alkyl;

R<sup>5</sup> is a hydrogen atom;

is 1; m

 $R^6$ is a phenyl; and is -COO-R12- $R^7$ 

wherein R12m is hydrogen atom, aralkyl, adamantyl, cyclohexylideneamino, piperidyl optionally substituted by lower alkyl, cyclohexyl optionally substituted by lower alkyl, or alkyl optionally substituted by a substituent selected from the group consisting of hydroxy, lower alkoxy, lower alkoxy lower alkoxy, lower alkoxycarbonyl, acyloxy, piperazinyl, and

amino optionally substituted by lower alkyl.

- 15. A pharmaceutical composition comprising a pharmaceutically acceptable carrier, and the amide compound of any one of claims 1 to 10 or a pharmaceutically acceptable acid addition salt thereof.
  - 16. An inflammatory cytokine production suppressor comprising the amide compound of any one of claims 1 to 10 or a pharmaceutically acceptable acid addition salt thereof as an active ingredient.
  - 17. An agent for the treatment or prophylaxis of an inflammatory diseases, comprising the amide compound of any one of claims 1 to 10 or a pharmaceutically acceptable acid addition salt thereof as an active ingredient.

## Amended claims under Art. 19.1 PCT

1. (Amended) An amide compound of the formula (I):

5	$R - A - X \stackrel{R^1}{\longleftarrow} M$	R <sup>2</sup> 0	(CH <sub>2</sub> )	R <sup>6</sup>	[)
	R <sup>3</sup>	R*	R <sup>⁵</sup>		

		R³	R*	R <sup>5</sup>
10	wherein;			
	R	is an optionally s R <sub>a</sub> , an alkoxy sub R <sub>a</sub> ,	ubstituted nor stituted by R <sub>a</sub> ,	a-aromatic heterocyclic group containing nitrogen, a hydroxy, an alkylthio substituted by $\mathrm{R}_{\mathrm{a}}$ , or an alkylamino substituted by
15		wherein R <sub>a</sub> is am carbonyl or imino group consisting protecting group;	these groups of lower alkyl	amidino, carbamoyl, ureido, thioureido, hydrazino, hydrazino- being optionally substituted by a substituent selected from the halogenated lower alkyl, cycloalkyl, aralkyl, aryl and amino-
	Α	is a linear or bra bond(s) in a chai		e which optionally has one or more double bond(s) or triple
20	v	is an exugen stor	n, or a single i	n, a cycloalkylene, a divalent aromatic heterocyclic group hav-
	X	ing one or more latom and oxyger CO-O-, -NH-CO-	hetero atom(s) n atom, -SO-, NH-, -NH-CS-	is selected from the group consisting of a nitrogen atom, sulfur -SO <sub>2</sub> -, -C=C-, -C=C-, -CO-, -COO-, -OOC-, -CS-, -COS-, -O-NH-, -NH-C(=NH)-NH-, -NR <sup>8</sup> -, -NR <sup>8</sup> CO-, -CONR <sup>8</sup> -, -NR <sup>8</sup> SO <sub>2</sub> -C-NR <sup>8</sup> -, or -CR <sup>9</sup> R <sup>10</sup> -
25	M	wherein R <sup>8</sup> is hyd and R <sup>10</sup> are the s is an arylene, a c	drogen atom, a same or differe sycloalkylene,	alkyl, cycloalkyl, aryl, aralkyl or amino-protecting group, and R <sup>9</sup> int and each is hydrogen atom, alkyl, cycloalkyl, aryl or aralkyl; or a divalent heterocyclic group which has one or more hetero
20		atom(s) selected and which option		p consisting of a nitrogen atom, sulfur atom and oxygen atom,
30	$R^1$ , $R^2$ , $R^3$ and $R^4$	are the same or of and R <sup>4</sup> is not a hy a nitro, a cyano,	different and ex ydrogen atom, a carboxy, an a	ach is a hydrogen atom provided that at least one of R <sup>1</sup> , R <sup>2</sup> , R <sup>3</sup> a hydroxy, a halogen atom, an alkoxy, a mercapto, an alkylthio, alkoxycarbonyl, an aryloxycarbonyl, an acyl, an alkyl optionally
35		halogen atom, as sisting of alkyl, a	n amino optior ryl, aralkyl and	lected from the group consisting of hydroxy, lower alkoxy and hally substituted by a substituent selected from the group condition of the substituted and obtained by a substituted of the substituted of t
40		cycloalkyl, option tuted alkylthio, o	nally substitute ptionally subs	ituted alkoxy, optionally substituted aryl, optionally substituted aryloxy, optionally substituted aralkyloxy, optionally substituted arylthio, or alkyl optionally substituted by a substituent sting of alkoxycarbonyl, acyloxy, aryl, aryloxy, aryloxycarbonyl,
	<b>8</b> ⁵	aralkyloxy, aralky	yloxycarbonyl, nally substitute	alkylthio, arylthio, acyl, lower alkoxy, carboxy, halogen atom ed by lower alkyl or acyl; optionally substituted by a halogen atom, an optionally substi-
	n	tuted aralkyl, or	an amino-prot	
45	m R <sup>6</sup>	is 0 or an integer is an optionally s	substituted ary	I, an optionally substituted cycloalkyl, an optionally substituted
50		amino optionally aryl, aralkyl and	substituted by amino-protect	ituted lower alkoxy, an optionally substituted lower alkylthio, an a substituent selected from the group consisting of lower alkyl, ing group, or an optionally substituted heterocyclic group haves selected from the group consisting of a nitrogen atom, sulfur
	R <sup>7</sup>	atom and oxyge	n atom; and	illy substituted alkyl, an optionally substituted aryl, an optionally
	n <sup>.</sup>	substituted aron	natic heterocyc	clic group having one ore more hetero atoms selected from the atom, sulfur atom and oxygen atom,
55		or -CO(Y) <sub>p</sub> R <sup>12</sup>	men atom eul	fur atom, -NR $^{13}$ - or -NR $^{13}$ -SO $_2$ -wherein R $^{13}$ is hydrogen atom,
		allow a railow by	drovy alkovy	aryl or amino-protecting group, p is 0 or 1, and R <sup>12</sup> is hydrogen
		atom ontionally	u cubetituted	elkenyl ontionally substituted alkynyl optionally substituted

atom, optionally substituted alkenyl, optionally substituted alkynyl, optionally substituted

cycloalkyl, optionally substituted aryl, optionally substituted aralkyl, adamantyl, cycloalkylideneamino, optionally substituted heterocyclic gr up having one or more hetero atom(s) selected from the group consisting of nitrogen atom, sulfur atom and oxygen atom, or alkyl optionally substituted by a substituent selected from the group consisting of hydroxy, alkoxy, alkoxyalkoxy, alkoxycarbonyl, acyloxy, carboxy, heterocyclic group having one or more hetero atom(s) selected from the group consisting of nitrogen atom, sulfur atom and oxygen atom, and amino optionally substituted by a substituent selected from the group consisting of alkyl, aryl, aralkyl and amino-protecting group;

10 or a pharmaceutically acceptable acid addition salt thereof.

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Α

Χ

M

R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup> and R<sup>4</sup>

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2. (Amended) The amide compound of claim 1, wherein, in the formula (I), at least one symbol selected from the group consisting of R, A, X, M, R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup>, R<sup>4</sup>, R<sup>5</sup>, m, R<sup>6</sup> and R<sup>7</sup> satisfies the following definitions, or a pharmaceutically acceptable acid addition salt thereof:

15 R is a non-aromatic heterocyclic group containing nitrogen, which is optionally substituted by lower alkyl or amino-protecting group, Ra1, an alkoxy substituted by Ra1, an alkylthio substituted by Rat, or an alkylamino substituted by Rat, wherein R<sub>81</sub> is amino, guanidino, amidino, carbamoyl, ureido, thioureido, hydrazino, hydrazinocarbonyl or imino, these groups being optionally substituted by a substituent selected from 20 the group consisting of lower alkyl, aralkyl and amino-protecting group;

> is a linear or branched alkylene which optionally has one or more double bond(s) or triple bond(s) in the chain, or a single bond;

is an oxygen atom, a sulfur atom, a cycloalkylene, a divalent aromatic heterocyclic group having one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, -SO-, -SO2-, -C=C-, -CE-, -CO-, -COO-, -OOC-, -CS-, -COS-, -O-CO-O-, -NH-CO-NH-, -NH-CS-NH-, -NH-C(=NH)-NH-, -NR8-, -NR8'CO-, -CONR8'-, -NR8'SO2-, -SO2NR8'-, -NR8'-COO-, -OOC-NR8'-, or -CR9'R10'-

wherein R8, is hydrogen atom, lower alkyl, aralkyl or amino-protecting group, and R9, and R10,

are the same or different and each is hydrogen atom, lower alkyl or aralkyl;

is an arylene, a cycloalkylene or a divalent heterocyclic group which has one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom,

and which optionally forms a fused ring;

are the same or different and each is a hydrogen atom provided that at least one of R1, R2, R3 and R4 is not a hydrogen atom, a hydroxy, a halogen atom, a lower alkoxy, a mercapto, a lower alkylthio, a nitro, a cyano, a carboxy, a lower alkoxycarbonyl, an aryloxycarbonyl, an acyl, a lower alkyl optionally substituted by a substituent selected from the group consisting of hydroxy, lower alkoxy and halogen atom, an amino optionally substituted by a substituent selected from the group consisting of lower alkyl, aralkyl and amino-protecting group, or

-O-CO-R111

wherein R11 is lower alkoxy, optionally substituted cycloalkyl, lower alkyl optionally substituted by a substituent selected from the group consisting of lower alkoxycarbonyl, acyloxy, aralkyloxy, aralkyloxycarbonyl, acyl, lower alkoxy, carboxy and amino optionally substituted by lower alkyl, or anyl optionally substituted by a substituent selected from the group consisting of lower alkyl, carboxy and benzyloxycarbonyl;

is a hydrogen atom, an alkyl optionally substituted by a halogen atom, an optionally substituted aralkyl, or an amino-protecting group;

is 0 or an integer of 1-6;

is an aryl, a cycloalkyl, or a heterocyclic group having one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom

wherein said aryl, cycloalkyl and heterocyclic group having one or more hetero atom(s) selected from the group consisting of nitrogen atom, sulfur atom and oxygen atom are optionally substituted by a substituent selected from the group consisting of lower alkyl, halogen

atom, hydroxy, lower alkoxy, amino, carboxy and lower alkoxycarbonyl; and

is a hydrogen atom, a lower alkyl optionally substituted by a substituent selected from the group consisting of hydroxy, lower alkoxy, mercapto, lower alkylthio, carboxy, lower alkoxycarbonyl and amino, an aromatic heterocyclic group which has one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, and

R7

R<sup>5</sup>

m R<sup>6</sup>

		which is optionally substituted by lower alkyl, or ${\rm -CO(Y')_p}{\rm R}^{12}$ , wherein Y' is oxygen atom, sulfur atom, ${\rm -NR}^{13}$ -, or ${\rm -NR}^{13}$ -, ${\rm SO}_2$ -wherein R <sup>13</sup> is hydrogen atom, lower alkyl, aralkyl, hydroxy, lower alkoxy or amino-protecting group,
5		p is 0 or 1, and R <sup>12</sup> is hydrogen atom, aralkyl, adamantyl, cycloalkylideneamino, cycloalkyl optionally substituted by lower alkyl, alkyl optionally substituted by a substituent selected from the group consisting of hydroxy, lower alkoxy, lower alkoxy lower alkoxy, lower alkoxycarbonyl, acyloxy, carboxy, heterocyclic group having one or more hetero atom(s) selected from the group consisting of, nitrogen atom, sulfur atom and oxygen atom, and amino optionally sub-
10		stituted by a substituent selected from the group consisting of lower alkyl, aralkyl and amino- protecting group, aryl optionally substituted by a substituent selected from the group consist- ing of lower alkyl, halogen atom, amino, carboxy, hydroxy and lower alkoxy, or heterocyclic group which is optionally substituted by a substituent selected from the group consisting of lower alkyl, halogen atom, amino, carboxy, hydroxy and lower alkoxy, and which has one or more hetero atom(s) selected from the group consisting of nitrogen atom, sulfur atom and
15		oxygen atom.
	group consisting of	amide compound of claim 1, wherein, in the formula (I), at least one symbol selected from the R, A, X, M, $R^1$ , $R^2$ , $R^3$ , $R^4$ , $R^5$ , m, $R^6$ and $R^7$ satisfies the following definitions, or a pharmae acid addition salt thereof:
20	R	is a non-aromatic heterocyclic group containing nitrogen, which is optionally substituted by lower alkyl or amino-protecting group, $R_{a2}$ , or an alkoxy substituted by $R_{a2}$ ,
		wherein $R_{\rm a2}$ is amino, guanidino, amidino or carbamoyl, these groups being optionally substituted by lower alkyl or amino-protecting group;
25	Α	is a linear alkylene or a single bond;
	Х	is an oxygen atom, a sulfur atom, a divalent aromatic heterocyclic group having one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, -COO-, -OOC-, -NR <sup>8</sup> "-, -NR <sup>8</sup> "CO-, -CONR <sup>8</sup> "-, -NR <sup>8</sup> "SO <sub>2</sub> -, -SO <sub>2</sub> NR <sup>8</sup> "-, or -CR <sup>9</sup> "R <sup>10</sup> "- wherein R <sup>8</sup> " is hydrogen atom, lower alkyl or amino-protecting group, and R <sup>9</sup> " and R <sup>10</sup> " are
30	М	the same or different and each is hydrogen atom or lower alkyl; is an arylene or a divalent heterocyclic group which has one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, and which option-
35	$R^1$ , $R^2$ , $R^3$ and $R^4$	ally forms a fused ring; are the same or different and each is a hydrogen atom provided that at least one of R <sup>1</sup> , R <sup>2</sup> , R <sup>3</sup> and R <sup>4</sup> is not a hydrogen atom, a hydroxy, a halogen atom, lower alkoxy, a lower alkyl optionally substituted by a substituent selected from the group consisting of hydroxy, lower alkoxy and halogen atom, or -O-CO-R <sup>11</sup> "
		wherein R <sup>11</sup> " is lower alkoxy, cycloalkyl, aryl optionally substituted by lower alkyl, or lower alkyl optionally substituted by a substituent selected from the group consisting of acyloxy,
40		aralkyloxycarbonyl and amino optionally substituted by lower alkyl;
	R⁵	is a hydrogen atom, a lower alkyl, or an amino-protecting group;
	m n6	is 1;
	R <sup>6</sup>	is an aryl or a cycloalkyl wherein said aryl and cycloalkyl are optionally substituted by halogen atom or hydroxy; and
45	R <sup>7</sup>	is a hydrogen atom, a lower alkyl optionally substituted by hydroxy or lower alkoxy, an aromatic heterocyclic group which has one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, and which is optionally substituted
		by lower alkyl, or $-CO(Y'')_pR^{12}$ " wherein $Y''$ is hydrogen atom, sulfur atom or $-NR^{13}$ "-wherein $R^{13}$ " is hydrogen atom, lower alkyl,
50		hydroxy or amino-protecting group, p is 0 or 1, and R <sup>12</sup> " is hydrogen atom, aralkyl, adamantyl, cycloalkylideneamino, cycloalkyl optionally substituted by lower alkyl, aryl optionally substituted by halogen atom, alkyl optionally substituted by a substituent selected from the group consisting of hydroxy, lower alkoxy, lower alkoxy, lower alkoxy, lower alkoxy, carboxy, heterocyclic group having
55		one or more hetero atom(s) selected from the group consisting of nitrogen atom, sulfur atom and oxygen atom, and amino optionally substituted by a substituent selected from the group consisting of lower alkyl, aralkyl and amino-protecting group, or heterocyclic group which is optionally substituted by lower alkyl, and which has one or more hetero atom(s) selected from

optionally substituted by lower alkyl, and which has one or more hetero atom(s) selected from

the group consisting of nitrogen atom, sulfur atom and oxygen atom.

4. (Amended) The amide compound of claim 1, wherein, in the formula (I), at least one symbol selected from the group consisting of R, A, X, M, R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup>, R<sup>4</sup>, R<sup>5</sup>, m, R<sup>6</sup> and R<sup>7</sup> satisfies the following definitions, or a pharmaceutically acceptable acid addition salt thereof:

R	is a piperazinyl optionally substituted by lower alkyl, a piperidyl optionally substituted by lower
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alkyl, an amino, or a lower alkoxy substituted by amino

wherein amino is optionally substituted by lower alkyl;

A is a linear alkylene;

X is an oxygen atom, a sulfur atom, -NH- or -CH<sub>2</sub>-;

M is an arylene;

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R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup> and R<sup>4</sup> are the same or different and each is a hydrogen atom provided that at least one of R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup>

and R4 is not a hydrogen atom, a hydroxy, a halogen atom, or -O-CO-R11"

wherein R<sup>11m</sup> is lower alkyl optionally substituted by a substituent selected from the group consisting of amino, acyloxy and benzyloxycarbonyl, or phenyl optionally substituted by lower

alkyi;

R<sup>5</sup> is a hydrogen atom;

m is 1;

20 R<sup>6</sup> is a phenyl; and

R<sup>7</sup> is -COO-R<sup>12</sup>···

wherein R<sup>12</sup>" is hydrogen atom, aralkyl, adamantyl, cyclohexylideneamino, cyclohexyl optionally substituted by lower alkyl, piperidyl optionally substituted by lower alkyl, or alkyl optionally substituted by a substituent selected from the group consisting of hydroxy, lower alkoxy, lower alkoxy lower alkoxy, lower alkoxy, piperazinyl and amino optionally substituted by lower alkyl.

- 5. The amide compound of claim 4, wherein M is phenylene, or a pharmaceutically acceptable acid addition salt thereof.
- 6. The amide compound of claim 4, wherein R<sup>7</sup> is -COO-R<sup>12</sup> wherein R<sup>12</sup> is lower alkyl, or cyclohexyl which is optionally substituted by lower alkyl, or a pharmaceutically acceptable acid addition salt thereof.
- 7. The amide compound of claim 4, wherein X is an oxygen atom or -CH<sub>2</sub>-, or a pharmaceutically acceptable acid addition salt thereof.
- 8. The amide compound of claim 4, wherein  $\mathbb{R}^6$  is phenyl and m is 1, or a pharmaceutically acceptable acid addition salt thereof.
- The amide compound of claim 4, wherein R is amino optionally substituted by lower alkyl, piperazinyl optionally substituted by lower alkyl, or piperidyl optionally substituted by lower alkyl, or a pharmaceutically acceptable acid addition salt thereof.
- 10. (Amended) The amide compound of claim 4, wherein R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup> and R<sup>4</sup> are the same or different and each is a hydrogen atom provided that at least one of R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup> and R<sup>4</sup> is not a hydrogen atom, hydroxy, a halogen atom, or -O-CO-R<sup>11</sup>"" wherein R<sup>11</sup>"" is lower alkyl or phenyl, or a pharmaceutically acceptable acid addition salt thereof.
- 11. (Amended) A carboxylic acid compound of the formula (I-a)

 $R \longrightarrow A \longrightarrow X \xrightarrow{R^1} \stackrel{R^2}{\longrightarrow} COOH$  (I-a)

wherein;

is an optionally substituted non-aromatic heterocyclic group containing nitrogen, a hydroxy, R  $R_a$ , an alkoxy substituted by  $R_a$ , an alkylthio substituted by  $R_a$ , or an alkylamino substituted by Ra, wherein R<sub>a</sub> is amino, guanidino, amidino, carbamoyl, ureido, thioureido, hydrazino, hydrazinocarbonyl or imino, these groups being optionally substituted by a substituent selected from the 5 group consisting of lower alkyl, halogenated lower alkyl, cycloalkyl, aralkyl, aryl and aminoprotecting group; is a linear or branched alkylene which optionally has one or more double bond(s) or triple Α bond(s) in the chain, or a single bond; is an oxygen atom, a sulfur atom, a cycloalkylene, a divalent aromatic heterocyclic group hav-10 X ing one ore more hetero atoms selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom. -SO-, -SO<sub>2</sub>-, -C=C-, -C=C-, -CO-, -COO-, -OOC-, -CS-, -COS-, -O-CO-O-, -NH-CO-NH-, -NH-CS-NH-, -NH-C(=NH)-NH-, -NR8-, -NR8CO-, -CONR8-, -NR8SO2-, -SO<sub>2</sub>NR8-, -NR8-COO-, -OOC-NR8-, or -CR9R10wherein R8 is hydrogen atom, alkyl, cycloalkyl, aryl, aralkyl or amino-protecting group, and R9 15 and R<sup>10</sup> are the same or different and each is hydrogen atom, alkyl, cycloalkyl, aryl or aralkyl; is an arylene, a cycloalkylene or a divalent heterocyclic group which has one ore more hetero M atoms selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, and which optionally forms a fused ring; and are the same or different and each is a hydrogen atom provided that at least one of R1, R2, R3 R1, R2, R3 and R4 20 and R4 is not a hydrogen atom, a hydroxy, a halogen atom, an alkoxy, a mercapto, an alkylthio, a nitro, a cyano, a carboxy, an alkoxycarbonyl, an aryloxycarbonyl, an acyl, an alkyl optionally substituted by a substituent selected from the group consisting of hydroxy, lower alkoxy and halogen atom, an amino optionally substituted by a substituent selected from the group consisting of alkyl, aryl, aralkyl and amino-protecting group, or -O-CO-R11 25 wherein R11 is optionally substituted alkoxy, optionally substituted aryl, optionally substituted cycloalkyl, optionally substituted aryloxy, optionally substituted aralkyloxy, optionally substituted alkylthio, optionally substituted arylthio, or alkyl optionally substituted by a substituent selected from the group consisting of alkoxycarbonyl, acyloxy, aryl, aryloxy, aryloxycarbonyl, aralkyloxy, aralkyloxycarbonyl, alkylthio, arylthio, acyl, lower alkoxy, carboxy, halogen atom 30 and amino optionally substituted by lower alkyl or acyl. 12. (Amended) The carboxylic acid compound of claim 11, wherein, in the formula (I-a), at least one of R, A, X, M, R1, R2, R3 and R4 satisfies the following definitions: 35 is a piperazinyl optionally substituted by lower alkyl, a piperidyl optionally substituted by lower R alkyl, an amino or a lower alkoxy substituted by amino wherein amino is optionally substituted by lower alkyl; is a linear alkylene; Α is an oxygen atom, a sulfur atom, -NH- or CH2-; Χ 40 M is an arylene; and are the same or different and each is a hydrogen atom provided that at least one of R1, R2, R3 R1, R2, R3 and R4 and R4 is not a hydrogen atom, a hydroxy, a halogen atom, or -O-CO-R11." wherein R11" is a lower alkyl optionally substituted by a substituent selected from the group consisting of amino, acyloxy and benzyloxycarbonyl, or phenyl optionally substituted by lower 45 alkyl. 13. (Amended) An amide compound of the formula (I-b) 50

5	x	is an oxygen atom, a sulfur atom, a cycloalkylene, a divalent aromatic heterocyclic group having one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, -SO-, -SO <sub>2</sub> -, -C=C-, -C=C-, -CO-, -COO-, -OCC-, -CS-, -COS-, -O-CO-0-, -NH-CO-NH-, -NH-CS-NH-, -NH-C(=NH)-NH-, -NR <sup>8</sup> -, -NR <sup>8</sup> CO-, -CONR <sup>8</sup> -, -NR <sup>8</sup> SO <sub>2</sub> -, -SO <sub>2</sub> NR <sup>8</sup> -, -NR <sup>8</sup> -COO-, -OOC-NR <sup>8</sup> - or -CR <sup>9</sup> R <sup>10</sup> - where n R <sup>8</sup> is hydrogen atom, alkyl, cycloalkyl, aryl, aralkyl or amino-protecting group, and R <sup>9</sup>
	М	and R <sup>10</sup> are the same or different and each is hydrogen atom, alkyl, cycloalkyl, aryl or aralkyl; is an arylene, cycloalkylene, or a divalent heterocyclic group which has one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom,
10	R <sup>1</sup> , R <sup>2</sup> , R <sup>3</sup> and R <sup>4</sup>	and which optionally forms a fused ring; are the same or different and each is a hydrogen atom provided that at least one of R <sup>1</sup> , R <sup>2</sup> , R <sup>3</sup> and R <sup>4</sup> is not a hydrogen atom, a hydroxy, a halogen atom, an alkoxy, a mercapto, an alkylthio,
15		a nitro, a cyano, a carboxy, an alkoxycarbonyl, an aryloxycarbonyl, an acyl, an alkyl optionally substituted by a substituent selected from the group consisting of hydroxy, lower alkoxy and halogen atom, an amino optionally substituted by a substituent selected from the group consisting of alkyl, aryl, aralkyl and amino-protecting group, or -O-CO-R <sup>11</sup> wherein R <sup>11</sup> is optionally substituted alkoxy, optionally substituted aryl, optionally substituted
20		cycloalkyl, optionally substituted aryloxy, optionally substituted by a substitutent selected from the group consisting of alkoxycarbonyl, acyloxy, aryloxy, aryloxycarbonyl, aralkyloxy, aralkyloxycarbonyl, alkylthio, arylthio, acyl, lower alkoxy, carboxy, halogen atom and amino optionally substituted by lower alkyl or acyl;
<i>2</i> 5	R <sup>5</sup>	is a hydrogen atom, an alkyl optionally substituted by a halogen atom, optionally substituted aralkyl, or an amino-protecting group; is 0 or an integer of 1-6;
	R <sup>6</sup>	is an optionally substituted aryl, an optionally substituted cycloalkyl, an optionally substituted lower alkyl, an optionally substituted lower alkoxy, an optionally substituted lower alkyl, an amino optionally substituted by a substituent selected from the group consisting of lower alkyl, aryl, aralkyl and amino-protecting group, or an optionally substituted heterocyclic group hav-
30	R <sup>7</sup>	ing one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom; and
35		is an optionally substituted aryl, an optionally substituted aromatic heterocyclic group having one or more hetero atom(s) selected from the group consisting of a nitrogen atom, sulfur atom and oxygen atom, or -CO(Y) <sub>p</sub> R <sup>12</sup> wherein Y is oxygen atom, -NR <sup>13</sup> - or -NR <sup>13</sup> -sO <sub>2</sub> -wherein R <sup>13</sup> is hydrogen atom, alled aralled bydrogen atom,
		alkyl, aralkyl, hydroxy, alkoxy, aryl or amino-protecting group, p is 0 or 1, and R <sup>12</sup> is hydrogen atom, optionally substituted alkenyl, optionally substituted alkynyl, optionally substituted cycloalkyl, optionally substituted aryl, optionally substituted aralkyl, adamantyl, cycloalkylideneamino, alkyl optionally substituted by a substituent selected from the group consisting of
40		hydroxy, alkoxy, alkoxyalkoxy, alkoxycarbonyl, acyloxy, carboxy, heterocyclic group having one or more hetero atom(s) selected from the group consisting of nitrogen atom, sulfur atom and oxygen atom, and amino optionally substituted by a substitutent selected from the group consisting of alkyl, aryl, aralkyl and amino-protecting group, or optionally substituted heterocyclic
45		group having one or more hetero atom(s) selected from the group consisting of nitrogen atom, sulfur atom and oxygen atom.
	14. (Amended) The the group consisting	amide compound of claim 13, wherein, in the formula (I-b), at least one symbol selected from of X, M, $\rm R^1$ , $\rm R^2$ , $\rm R^3$ , $\rm R^4$ , $\rm R^5$ , m, $\rm R^6$ and $\rm R^7$ satisfies the following definitions:
50	X M	is an oxygen atom, a sulfur atom or -NH-; is an arylene;
	R <sup>1</sup> , R <sup>2</sup> , R <sup>3</sup> and R <sup>4</sup>	are the same or different and each is a hydrogen atom provided that at least one of R <sup>1</sup> , R <sup>2</sup> , R <sup>3</sup> and R <sup>4</sup> is not a hydrogen atom, a hydroxy, a halogen atom, or -O-CO-R <sup>11</sup> wherein R <sup>11m</sup> is lower alkyl optionally substituted by a substituent selected from the group
55	-5	consisting of amino, acyloxy and benzyloxycarbonyl, or a phenyl optionally substituted by lower alkyl;
	R <sup>5</sup> m	is a hydrogen atom; is 1;

		El O OTO BOO AT
5	R <sup>6</sup> R <sup>7</sup>	is a phenyl; and is -COO-R <sup>12</sup> ···· wherein R <sup>12</sup> ···· is hydrogen atom, aralkyl, adamantyl, cyclohexylideneamino, piperidyl optionally substituted by lower alkyl, cyclohexyl optionally substituted by lower alkyl, or alkyl optionally substituted by a substituent selected from the group consisting of hydroxy, lower alkoxy, lower alkoxy, lower alkoxy, piperazinyl, and amino optionally substituted by lower alkyl.
10	any one of claims 1	tal composition comprising a pharmaceutically acceptable carrier, and the amide compound of to 10 or a pharmaceutically acceptable acid addition salt thereof.
	or a pharmaceutica	y cytokine production suppressor comprising the amide compound of any one of claims 1 to 10 lly acceptable acid addition salt thereof as an active ingredient.
15	17. An agent for the one of claims 1 to 1	treatment or prophylaxis of an inflammatory diseases, comprising the amide compound of any 0 or a pharmaceutically acceptable acid addition salt thereof as an active ingredient.
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### INTERNATIONAL SEARCH REPORT International application No. PCT/JP96/02305 CLASSIFICATION OF SUBJECT MATTER Int. C16 C07C235/60, 279/ C07C235/60, 279/08, C07D211/34, 241/04, 295/08, 295/10, 263/58, 271/06, A61K31/215, 31/445, 31/495 According to International Patent Classification (IPC) or to both national classification and IPC FIFI DS SEARCHED Minimum documentation searched (classification system followed by classification symbols) Int. C1<sup>6</sup> C07C235/60, 279/08, C07D211/34, 241/04, 295/08, 295/10, 263/58, 271/06, A61K31/215, 31/445, 31/495 Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) CAS ONLINE C. DOCUMENTS CONSIDERED TO BE RELEVANT Category\* Citation of document, with indication, where appropriate, of the relevant passages JP, 63-238051, A (Showa Denko K.K., Yusuke 1-3, 11, 12, Х 15, 17 Okamoto), October 4, 1988 (04. 10. 88), 4-10, 16 Claim; pages 5 to 11 (Family: none) Α JP, 63-239256, A (Showa Denko K.K., Yusuke 1-3, 11, 12, Х Okamoto), 15, 17 October 5, 1988 (05. 10. 88) Claim; pages 6 to 16, 20 to 22 (Family: none) 4-10, 16 Α 1-3, 13, 15, JP, 48-18241, A (Imperial Chemical Industries Х Ltd.), 17 March 7, 1973 (07. 03. 73), Claim; page 2; page 5, lower left column; page 8, upper left column & GB, 1391444, A 14, 16 Α & CH, 573393, A & CH, 575908, A £. See patent family annex. Further documents are listed in the continuation of Box C. later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention Special categories of cited documents: document defining the general state of the art which is not considered to be of particular relevance "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "E" earlier document but published on or after the international filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art "O" document referring to an oral disclosure, use, exhibition or other document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of mailing of the international search report Date of the actual completion of the international search November 7, 1996 (07. 11. 96) November 19, 1996 (19. 11. 96) Name and mailing address of the ISA/ Authorized officer Japanese Patent Office Telephone No.

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